# A SEROEPI DEMIOLOGY OF MALARIA IN A RURAL POPULATION OF PRIMARY HEALTH CENTRE, CHIRGAON, JHANSI (U.P.)

# THESIS DOCTOR OF MEDICINE

(SOCIAL AND PREVENTIVE MEDICINE)



BUNDELKHAND UNIVERS
JHANSI (U. P.)



# CERTIFICATE

This is to certify that the present work entitled "A SERO-EPIDEMIOLOGY OF MALARIA IN RURAL POPULATION OF PRIMARY HEALTH CENTRE, CHIRGAON, JNAMSI (U.F.) " has been carried out by Dr. KAMAL KISHORE REMY himself in this department.

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# CERTIFICATE

This is to certify that the present work entitled "A SERO -EPIDEMIOLOGY OF MALARIA IN A KURA POPULATION OF PRIMARY HEALTH CENTRE, CHIRDACK, JHANSI (U.P.)", has been carried out by Dr. KAMAL KIGHORS REMY, under our constant supervision and cuidence. The observations were checked and verified by us from time to time.

This thesis fulfile the basic ordinance governing the submission of thesis for M.D., laid down by Bundelkhand University.

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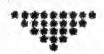
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INTRODUCTION

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#### INTRODUCTION

Malaria has reappeared during the last decade throughout the tropical and subtropical regions of the world. In the world today, there have been radical changes in socio-economic conditions and some consequences of development have led to changes in habitats and acosystems of malaria vectors and parasites. Developments in agriculture such as intensification and reorientation, green revolution, irrigation, deforestation, development in the industry and other technical interventions; high mobility of population as well as increased labour force movement, resettlement and rehabilitation, increased tourism, have all introduced new aspects in the epidemiology of malaria. The development of drug resistance by the parasite and insecticide resistance by the vectors are the prime causes for the reappearance of malaria in the areas where it was practically controlled previously. Veilure of operational and surveillance measures were also equally responsible for the resurgence of malaria.

The revised global plan for achieving the containment of the virulent malaria with the ultimate aim of eradication of the disease envisages, among many

other aspects, a proper estimation of the extent to which population in endemic creas has been exposed to malaria.

it is observed that malaria had, in fact, been erradicated in over 20 countries, freeing a population of nearly 800 million from risk of the disease and transmission of the disease had also been gradually reduced in areas inhabitated by 775 million people.

It has also been observed that modification of clinical illness occurs in endesic areas, due to development of immunity. In such areas parasitaexia way be seen without clinical illness and vice versa and also the individuals may be missed by screening of fever cases only. Parasitaenia in malaria is intermittent and its absence on single slide examination does not exclude the diagnosis of malaria. This limits the usefulness of slide examination for malaria detection epidemiologically. Secondly, slide surveys of the population are cumbersome and do not yield results commensurate with the work involved. Another indicator of prevalence of melaria has been the splean rate. This is a good index for giving on the spot discnosis, but it does not hold good in areas with low endesicity. Resides malaria, there are other causes of splenomegaly and it is not enlarged in all patients of malaria, hence there is

a need for a newer method, which is better indicator of endemicity and transmission of malaria and involves less expense, time and labour.

reported to be useful tool for studying malaria endemicity rates, patterns of malaria transmission and to detect foci of malaria in epidemiological survey (kagar, 1972).

Of large number of serological tests available,
ELISA & IIF test have been found to be simple, reliable,
reproducible sensitive, specific and large number of
samples can be economically processed (Voller et al, 1980).

Many studies have been conducted abroad but it needs
evaluation is Indian conditions. Another aspect of
serology of malaria which needs evaluation, is the use
of serology for the diagnosis of individual patient or
community diagnosis. No single test has been found
useful in this respect but use of sultiple tests needs
evaluation. Resping these facts in mind, the present
study has been designed with the following sime and
objectives:

 To assess the prevalence of antibody titre in random population.

- To find out correlation between sero-positivity and alide positivity.
- 3. To evaluate application of serology to study the epidemiology of malaria by correlating it with various bio-social characteristics of the population.

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REVIEW OF LITERATURE

#### REVIEW OF LITERATURE

The word malaria is derived from an Italian word, Mala-aria which stands for bad airs. In 1740, Morace Walpole, for the first time used the word Malaria in English.

The term Kalaria is applied to a group of diseases caused by infection with specific sporosoon parasites of genus plasmodium and transmitted to man by various species of Anopheline mosquito transmit the disease which is clinically characterised by episodes of chills and fever with period of latency, emlargement of spleen and secondary emacmic (Park & Park, 1989).

Malaria is caused by four distinct species of plasmodium, viz. P. vivax, P. falciparum, P. malariae and P. ovale. Agent requires two hosts for its propagation and completion of life cycle. The disease affects individuals of all ages and both sexes especially of low socio-economic status, living in ill-ventilated, ill-lighted and unhygienic houses surrounded by various types of water collections. The female anopheline mosquito, the definitive host, requires 20 - 30°C temperature and high humidity for active life. Such conditions prevail during rainy season, when the disease flares up.

Though the disease is wide-spread all over the tropical belt. it presents with varying degree of endemicity. the seasonal variation host and vector factors responsible for the disease. There is need for an economical, simple, safe, reliable, sensitive, specific method for assessing the extent of the problem. Three methods viz. spleen rate, perasite rate and serology are in use. Ideally, a method should identify the infecting species differentiate between present and past infections and indicate the immune status of the host. A test that fulfilled these goals would be both sensitive and discrestic sethod and also a valuable tool for the study of epidemiology of maleria. In view of this, serology seems promising in areas of high incidence malaria; the age groups above 4 years acquire immunity induced by repeated infections, resulting more number of unrecognized cases of malaria.

Although no method has fulfilled these criteria; yet some serological methods have proved to be a valuable tool to assess more accurate appreciation of the prevalence of malaria, especially in circumstances where the use of enti-malarials, invalidates the classical parameters of malaria endemicity such as the prevalence of spleen enlargement and parasitaemia (N.H.O., 1975).

# 1. Mistory of Disease :

Ancient history during vedic times, malaria is reported as "King of diseases" and was often attributed

due to anger of lord Shive in medical literature (Charaka Samhita & Susrute Samhita, 600 B.C.).

The Chinese, centuries before the Christian era. differentiated tertian from quertan fewers and recognised the enlargement of spleen in selects.

The year 1880 is important in the history of malaria as it marked the discovery of malaria parasite in fresh human blood by Laveran and then Romanowski gave to the world an original technique of staining blood smears.

Ronald Ross (1898) discovered the life cycle of Malaria parasite in the invertebrate host (The mosquito). The introduction of plasmochin (now called Pamaquin, 1924). a synthetic anti-malarial in its chamotherapy followed by Atebrine (1930) now called mepacrine, chloroquin (1934) and Faludrine (Froquanil, 1945) were outstanding discoveries in the field of malariology (Central Health Education Bureau, C.G.H.S. Ministry of Health, Gowt. of India).

Piratly, in the control methods re-discovery of

Pyrethrin (1936) a contact insecticide marked the beginning

of maleria control measures and later, followed by residual

synthetic insecticides like D.D.T., B.H.C., Dieldrin and

other organophosphorous compounds etc. and proved practicable.

Sconomically feasible in rural areas, towns and cities

(Central Health Education Bureau, 1959).

Morld Health Organization from its inception has recognised malaria as a major health problem, out of six tropical diseases, malaria was the target of N.H.J. special programme for research and training in tropical diseases to develop new tools and strengthen research institutions and training workers in the countries affected (Park & Park, 1989).

# 2. Epidemiological features of Contemporary Malaria :

Malaria is widely prevalent throughout the world with a high prevalence in Asia and Africa. It was established that the population at risk of malaria (excluding China), was about 1729.17 million in 106 countries of the world. Out of which, the eight countries of South-East Asia contributed for 918.72 million people (N.H.O., 1962).

the resurgence of salaria had occurred throughout the world, with a peak in 1976 followed by decrease in the number of cases so that by 1982 the number was almost similar to that reported in 1974 (TRS - 735). By end of 1974, there were 1,136 million people at the risk to suffer from malaria (Srivestava et al. 1975) in India.

# 2.1 Demographic explosion :-

The average growth of more than 2% population was significantly increased among economically backward

classes, inhabiting areas with difficult accessibility on the periphery (Ray, 1979; Kondra Shin, 1983).

# 2.2 Intensification and Re-orientation of agriculture :

About 70 to 80 percent of population being engaged in ferming in the country. It was estimated that there were about \$8 million people engaged in seasonal egricultural activities and moving for purposes of harvesting throughout the country (Kondrashin, 1983).

# 2.3 Industrial Development :

production of steel. electronic goods, heavy machines, fertilizers, exploration of oils, bamboo cutting, lend clearance, gem-mining, coal fields, ore mines, thermal power stations, roads, lime store, dolomity, quarts, aluminium, copper etc. The mineral wealth of the country lies in areas of difficult terrain, mostly in hills, which are hard core areas (Ray, 1979).

# 2.4 <u>Urbanization</u>:

It may be seen that both natural (high temperature and humidity vast areas of vector breeding places, prolonged transmission season etc.) and socio-economic factors (low sanitary standards, intensive population movement, demographic explosion etc.) create extremely favourable conditions for transmission of vector horne diseases in urban areas.

#### 2.5 Improvement in transport facilities :

projects, the road communication has increased tremendously. It is estimated that around 50% of labour is imparted from other places in the country consisting of landless workers. They carry infection from hard core areas to urban as well as rural areas.

#### 2.6 <u>Deforestation</u> :

In forest based industries, shifting agriculture is still in use, resulting in massive deforestation which in turn results in degradation of land, soil erosion and sedimentation of lakes, river and reservoirs with increase in the density of mosquito (Kondrashin, 1987).

# 2.7 Irrigation :

On the whole, it appears that in India, canal irrigation leads to a marked rise in the ground water table and any disturbance in natural drainage contours sometimes obstructs the natural drainage flow of rain water in the area, such changes create favourable malariogenic condition with high transmission potential.

# 2.8 Unemployment & under-employment :

The most important of Indian economy, the unemployment problem is essentially rural in character and has grown vastly in magnitude the absolute number of unemployed on a typical day about 21 million constituting 8.2 percent of the labour force (Agarwal, 1985). There could be an epidemic out-break of malaria in the labour camps and increased vulnerability of malaria in the labour camps and in the originating areas on account of the return of labourers.

The labour force are being made at a fast rate on account of the rapidly growing population. Thus, while new productive jobs are on the increase, because of the low rate of increase, the absolute number of unemployed persons is rising from year to year (Agarwal, 1985).

# 2.9 Tourism and Pilgrimage :

The celebration of religious festivals like Numbha Hela results in the congregation of many hundreds of thousands even millions of people at a time. The results of random study conducted on a group of pilgrims for several states of India have shown that the parasite rate among them was about 2% and all cases were due to P. felciparum. a symptomatic gametocite carriers (Raj Gopelen et al. 1986).

Malaria is not only a rural problem but is also important in urben areas.

# 3. Ace/Sex/Occupation related Malaria :

# 3.1 Age related malaria :

The prevalence of malaria among different age groups is subject to wide variations. In a primary health centre Mainital district of Utter Fradesh, Choudhary et al (1984) carried a study and classified the population on the basis of clinical history of malaria observing that all age groups were affected by disease but that there was a progressive increase of malaria attacks from infents to 15 - 25 years of age, when the rates reached the maximum level the NICD has conducted serological studies in hypo-endemic malarious areas of the country during the 1970s and it was observed that the population below 5 years of age had hardly any malaria experience. It was only higher age groups who showed high titres. Kummar et al (1986) in a study around Delhi i.e. Somepaty Gurgaon and Gaziabed, revealed ELISA antibodies showed a very definite trend towards age related increase in the titre, employing P. falciparum antigen, gave desired results. Both the IMA & ELISA meen titres correlate well with S.P.R. in the non-transmission period such a correlation was lacking in other period, the expected age related increase in antibody titres was evident with ELISA and not DIA.

Rumer et al (1987) reported that the immunological profile of a population was the sum total of previous individual experiences. The response of the individual

to these experiences was affected by age, immunological competence, cumulative exposure to malaria antigen and kind and amount of specific therapy used, and also reported age related increase in the number of individuals with positive peripheral blood smears increase upto the age of 4 years and then there was a sharp decline. The infection rate was found to be 26.9 in the age group 1-5 years. The infection rate was calculated for each willage individually which ranged from 1.7 to 63.2. There was a good correlation in the age group 1-4 years between the infection rates of each village and total number of malaria cases as percentage of entire population of each village.

# 3.2 Sem related melarie :

Sex differences between the general distribution of these in Indian population and in the percentage of malaria cases reported for each category probably are influenced by local socio-economic status, ethnic groups, the attitude of parents especially mothers towards males and female children treatment for malaria, ignorance about the availability of free services in the village. In Utter Pradesh state, Chowdhary et al (1984) observed that both sexes were affected by the disease but incidence among males was almost twice as high as compared with that among females.

# 3.3 Occupation related malaria :

Different occupational categories of labour force were identified as being at increased risk to acquire malaria, including its resistant forms and disseminating the disease. The overwhelming majority of total malaria cases annually occurs among various categories of agricultural labour (Pattanayak, 1981). The rest of the cases occur in urban and other areas of the country (Sharma, 1984; Kondrashin and Dixit, 1985). The risk to acquire malaria is high among mobile workers and among those exposed to mosquite bites in the open air on account of their occupational requirements (Kondrashin, 1986). The higher S.P.R. was reported in labourers engaged in bamboo cutting, tea plantations, coal fields, coconut plantation, fishermen (Panicker et al., 1984; Panicker & Rajgopalan, 1986).

# 4. Malaria in India :

Malaria is one of the great scourages afflicted humanity. Even early in this century there was no aspect of life in our country which was not affected either directly or indirectly by Malaria (Sinton, 1936).

At the time of independence, malaria was regarded as a major public health problem. The annual incidence of malaria was 75 million cases with 8 lakh deaths directly due to malaria (Park 6 Park, 1989). In post independent era,

Gowt. of India realizing gravity of problem, started National Malaria Control Programme in 1953, which upgraded to National Malaria Eradication Programme in 1958 due to fear that vector may develop resistence to insecticides (Malaria in India, 1958).

of malaria dropped down from 75 million cases to 2 million cases in 1958 and the proportional case rate fell from 10.8 in 1953 to 3.2 in 1958 (Park & Park, 1989). Progress in malaria control was also seen in neighbouring countries of South East Asia Region of W.H.O. Five had progressed beyond the attack phase except Nepal, 19 - 77% of population of other five countries had reached the 'consolidation phase'.

In India itself, 99% population was covered by attacked phase and 1% by consolidation phase in 1961.

By 1966, 14% population was in attack phase, 34% in consolidation phase and 52% was under maintenance phase.

The success of the malaria eradication programme was due to normal sensitivity of parasite to chloroquin, and sensitivity of anopheline vectors to D.D.T. However, programme suffered a set back after 1965 in the form of small and large focal outbreak of malaria in different states of the country. Disease showed an upsurge trend year after year till the condition started worsening from 1975. There were 13,58,753 cames of malaria in 1971 which

increased to 53,10,790 in 1975. Considering the gravity of the situation, a modified plan of operation was implemented in 1978.

There were 64,67,215 cases of malaria in 1976 which decreased to 17,65,631 in 1986. The status of malaria in the country in last 11 years is shown in table 2.1.

Table 2.1

70.83	Total malaria cases	P.falciparum cases	ARRUAL parasite incidence	8.2.8.
1976	64.67,215	7,53,713	11.24	11.553
1977	47,40,900	4,59,867	8.07	8.316
1978	41,44,385	5,48,567	6.80	6.855
1979	30,64,697	5, 58, 423	4.90	4.990
1980	28, 98, 140	5, 98,011	4.51	4.315
1981	27,01,141	5, 89, 591	4.11	3.982
1982	21,82,302	5, 51, 057	3.22	3,356
1983	20, 18, 605	6,00,694	2.93	3.140
1984	21,84,446	5, 86, 691	3.00	3.290
1985	18,64,380	5,45,005	2.57	2.740
986	17,65,631	6,21,235	2.40	2,660

Source : W.H.O./SEARO, 1987.

Melaria is not only a rural problem but is also important in urban areas. Fatteneyak gi al (1981) had shown that Madras city contributed 40 - 50% of total malaria cases of the state. The number of malaria positive cases in ten major cities in India between 1978 to 1985 are shown in Table 2.2.

Table 2.2 Nalaria cases in ten major cities in India (1978, 1980-1985).

	Fop. in	and the later of t	at an affiliance of a first first from the latest	Mala	cia ca	100		
Cities	sillion	1978	1980	1901	1982			
Ahmedabed	2.52	26705	26664	20703	12006	18329	23186	16113
boahay	8.23	2635	1608	309	F.A.	3700	2610	1371
taroda	0.74	29866	12161	13648	8743	8378	5769	6562
lengelore	2.91	952	322	216	101	59	34	31
thop al	0.67	2656	2893	H.A.	1103	2055	2740	2746
heedigarh	0.42	34748	36278	31209	25945	28835	24035	36543
alcutta	9.17	1244	3246	5527	5304	19370	26056	21303
Sellei	5.71	332683	69277	62415	46530	4107	38108	28577
lyderabed	2.53	2559	1242	2494	4337	2400	3346	4096
edras	4.28	24953	36193	43981	44981	44817	48523	51376

Source : MMEP, Delhi 1987, based on 1981 census.

Table 2.2 shows that the cities of Delhi, Madras,
Calcutta, Ahmedebad and Chandigarh are the main contributors
to the problem of urban melaria in India at present. This was

particularly so in the year 1978, when the number of maleria cases in only ten major cities of the country accounted for more than 11 percent of total malaria cases in the country. This problem have infact been on the increase since the year 1982 onwards.

In Lakshadweep Island, Ray et al (1978) observed the A.F.I. of 102, 445.2, 20.2 from Mirucoy, Bitra and Chatlet Islands respectively. In all above studies parasite was sensitive to chloroquin.

Shanmugham et al (1978) from Tamil Nadu also reported 6656 cases of <u>P. Palciperum</u> and treated them with 1200 mg and 600 mg chloroquin base for adults and children respectively. Monthly follow-up of these patients was done after the treatment. Daly one patient showed parasite in peripheral blood. Soon it was observed that P. Falcidarum in certain parts of India had developed chloroquin and sultiple drug resistence. The chloroquine resistance against F. Falciparum was first detected in Assam (Sengal et al 1973) subsequently chloroquine resistance was detected in Arunachal Pradesh, Misorum, Mechalaya and Magaland (Pattanayak et al 1979; Chakraborty et al 1979 and Das et al 1979). The resistence was also found in Maharashtra, Orissa, Ottar Pradesh and Madhya Pradesh (De, et al. 1979; Suha et al. 1979 and Dwiwedi et al. 1981; Baveja et al. 1985). These strains are apreading in wirelent fore densing high degree of morbidity and mortality in children. Krotochi(1981)

observed that some South East Asian strains of  $\underline{P}_{\bullet}$  yives were also resistant to standard regimens of Privaquin.

Melaria control in rural areas was carried out by spraying residual insecticides such as DDT, HCH or malathiam. In areas with DDT resistant vectors, HCH was sprayed. At present about 210 million population is under DDT spray and about 100 million under HCH spray.

Malathion resistance in A. culicifacies developed in areas of Orissa and Andhra Fradesh, where this insecticide was never used in public health, this development of resistance in A. culicifacies was the result of the use of organo phosphate compounds in agriculture (Magpal, 1986; Sharma, 1987 b). At Shahjahanpur, the bio-environmental control of malaria strategy was implemented in 1986, results so far achieved have shown major reduction in vector densities and SPR was reduced from 80 - 90% to 20 - 30% (Sharma, 1987 b).

# 5. Malaria in Utter Prodesh :

Utter Predesh is the largest state of India with a population of 110.9 million (Census, 1981).

Meleria eradication programme was launched in the state during 1958-59. By 1959-60, sixty seven eradication programme units were established in the state covering 67 million population of the state. Jerring the population

living in areas above 5000 ft, the entire state was covered under the programme (State Health Education Fureau, U.F., 1987).

The entire state of Uttar Predesh remained in attack phase till 1961-62. The Units which had achieved the desired criteria were recommended for entry into enother phase by the independent Appraisal Teams, Started entering into consolidation phase' from 1962-63 and into maintenance phase from 1965-66. By 1969-70, out of 67 units, \$1.27% units entered into maintenance phase (State Health Education Bureau, U.P., 1987).

During the period 1951-1964 the incidence of salaria was brought down to a very low level and the state was almost free from malaria. The programme suffered set backs in some units from 1965, and the disease spreaded and showed an upward trend year after year from 1965 to 1977. 4,33,944 cases were seen in 1977. The year-wise data from 1970 to 1977 is as follows (Table 2.3).

Table 2.3

Year		Wo.of	malaria	Cases	detected
1970				002	
1971			9,	798	
1972			17,	199	
1973			54,	145	
1974			1,90,	755	
1975			3,81,	750	
1976			3,87.	728	
1977			4,33,1	744	

There has been a considerable improvement in equidemiological situation since 1977. But during 1979 the incidence of malaria cases was on the increase till 1984. However, during 1985 the incidence of malaria has declined by 11% as compared to the year 1984. The incidence of malaria cases was further declined 66.17% in the year 1987 as compared to 1985 (Table 3) (State health Education Bureau, U.P., 1987).

Table 2.4

Epidemiological data from the years 1977 onwards is as follows:

****	Positives	Astal	5.2.2
1977	4,33,944	4.4	5.8
1978	3,60,059	3.3	4.3
1979	1,49,919	1.5	1.9
1980	1, 82, 308	1.7	1.9
1401	1,75,930	1.6	1.9
982	1,70,233	1-6	2.2
1903	2, 85, 618	2.6	3.5
1984	4,19,708	3.6	4.5
1985	3,73,006	3.1	4.2
1986	2,28,244	1.09	2.9
1987	1,26,181	1.04	1.50

API : Annual parasite incidence, SPR : Slide positivity rate. Source : Deptt. of halariology. Govt. of Utter Predesh.

# 6. Kalerie in Jhansi :

It has the population of 11,37,031 and having 936 villages (Census, 1981). This district is having hot and dry climate. Srivastava et al (1975) in a study had showed the mean monthly maximum and minimum temperature range between 24.1°C to 42.6°C and 9.2°C to 29.3°C respectively. The mean monthly relatively humidity ranges between 26 to 84% at 0830 hrs. and 15 to 76% at 1730 hrs. Mean monthly rainfall ranges from 2.7 mm in the month of April and 139.1 mm. in the month of August. There were 104 surveillance units with a total of 1312430 surveillance population in 1973 (Szivastava et al. 1975). Data presented in the table 2.5 reveals that the incidence of malaria cases showed a rising trend from 1980 to 1984, thereafter they started declining. It was observed that there was a fall of about 92% in the number of positive cases in the year 1987 in comparison to 1984.

Table 2.5

Year	Incidence of maleria in Jhansi	A.P.I.
1980	1434	7.6
1981	2155	6.7
1982	1869	7.0
1983	2093	7.0
1994	7594	
1985	7591	5.6
19 36	5444	5.1
1987	3991	3.7

Source : District Melaria Office, Jhansi.

In the year 1987, status of malaria at Frimary Mgelth Centres of Jhansi is shown in table 2.6.

Table 2.6

Name of P.H.C.	Ro. of melaria cases detected	Askala	S.F.A.
Patina	555	4.9	7.4
Meuranipur	682	4.7	7.5
Ture ared	76	0.64	0.75
leas-	159	1.4	2.5
Sarageon	371	3.6	9.6
thirg aon	275	2.6	2.8
woth	650	4.7	6.8

Source : District Malaria Office, Jhansi.

In a study, Srivastava st sl (1975) and Verma

st sl (1975) had found that P. vivaz infection was

responsible for 92.25% morbidity and A. culicifacies and

A. fluviatilis are two known vector of malaria in Jhamsi,

with former playing the major role in the transmission of

the disease. They have also observed favourable

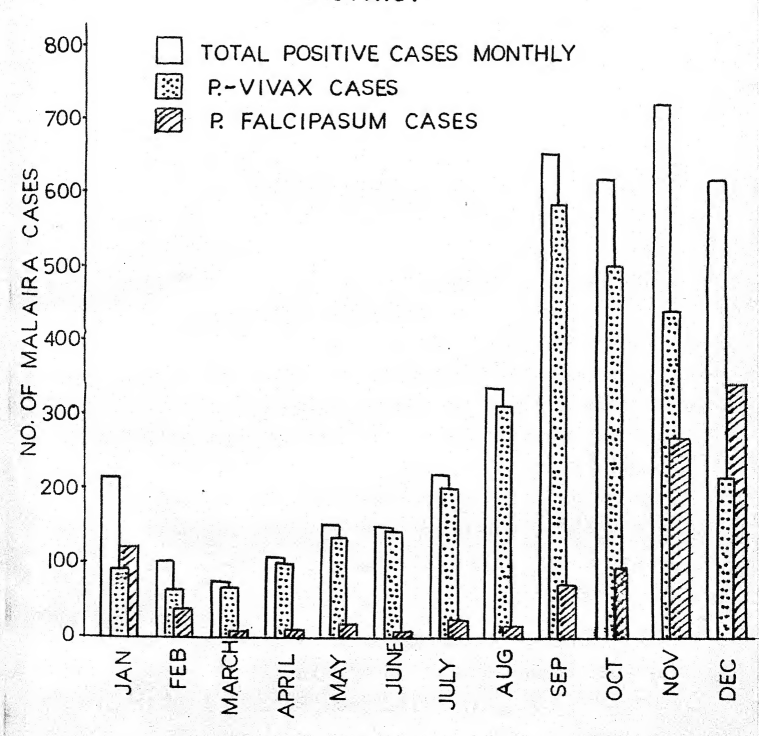
transmission season for the species P. vivaz and P.facliparum.

from August to October as peak month and minimum load of

infection during January and December. The monthly malaria

positive cases in Jhamsi depicted in figure 2.1.

FIG - 21
BAR DIAGRAM SHOWING MONTHLY MALAIRA
POSITIVE CASES DURING 1987 IN JHANSI
DISTRICT



# 7. Spleen rate :

The spleen rate is the proportion of children (sgad 2 - 10 years) in a community who have large apleen. The apleen rate includes past as well as present malaria infection and it is fracquently more reliable in practice than examination of blood films (Mac Donald, 1957). Mackett's method is used for classification of enlarged splean. Splean size is not considered for determining the spleen rate itself, but it is used to calculate average enlargement of spleen, another indicator of endemicity. In the holo- or hyper-endemic areas of melaria, the A.S.S. in ace group 2 - 9 years is high and in the epidemic areas is low in the same age group. After interruption of malaria transmission in highly endemic areas, splean rate remain the same for months or diminish insignificantly but on the other hand the A.E.S. remarkably decreases, therefore, is a more sensitive tool in evaluation than the spleen rate alone (Marshal, 1986). The spleen rate taken among the 2 - 9 years age group and determination of A.E.S. are easier to obtain than parasite rate.

Spleen rate is a good index for diagnosis of malaria during epidemics and in hyperwendemic areas as it gives on the spot epidemiological situation, the degree of immunity, previous history and future prospects of disease. No other disease is known to cause such a high

spleen rate except Kala Azar. But then, all patients with splenomegaly do not have malaria and all patients of malaria do not have splenomegaly.

#### 8. Parasite Rate :

The parasite rate is the proportion of a population in which malaria parasites are found. Infant parasite rate is defined as the percentage of infants below the age of one year showing malaria parasites in their blood. However, it is an important procedure in both individual diagnosis and epidemiology and has been main stay of parasitic detection in infected mosquitoes and in man, till relatively simple diagnostic tests are developed (N.H.O., 1987).

Blood examination for malaria parasites provide a seasonably adequate measure of the point prevalence of the infection. For community application, it requires large number of field workers as well as technicians.

Fon-fever cases are not studied, so sub-clinical infection without fever remains undetected. In a study in meao-endemic area, Upreti et al (1982) obtained 6.8% slide positivity rate in a febrile healthy children of 2-9 years, although the positivity was higher (45.5%) in febrile ones. Slide examination can only indicate the presence and absence of patient parasitaemia at the time of examination; it does not indicate the individuals malaria experience (Kagam, 1972).

The results of spinen surveys are compared with other epidemiclogical methods in following table. Comparison of methods of epidemiological investigations in malaria.

- 6		Spiesa rate X	Paraelte rate %	ATATATROA	******
:	Sherma at pl (1965) (2-9 years)	30.65	3. 3.	Kot dobe	Splean was found in many parasitolo- gically negative cases.
*	Upreti at al (1962) (2-9 years)	****			splenomegaly detected in meny parasito-
	Mamteni et al (1979)		7.0	36.0	
	7/ 15 yrs.	10.0	37.0	•	Results of mero-positivity and parasite rate provide a reasonably adequate measure of point prevalence and degree of malaria endemicity.
	Xeay 25 al (1973)	0.04			There was no correlation between splenomedaly and sercobaltivity.
;	Colling & Millian  1: 1, (1972)  1: 5 years		0	***	3
	Area (b) 1-5 years	Not con	18.0	9 9	iffering picture of
	7 15 years		22.6	***	
	Descuits at al (1965)	0.5	72.2		
•	2-9 yearst (1) (1) (2)	44 60 60	6.50	000	Spleen rate was much lower than parasite rate and seropositivity.
	S (1941) 42 42 1.944 (8)	meo *****	***	Not done	In low endemic areas spleen rate is not useful.

The absence of patent parasitaemia can be misleading since patency is influenced by immune status, and use of anti-malarial drugs. Besides parasites may not be present in peripheral blood continuously during cycle, especially in falciparum malaria where only ring stages develop in peripheral blood and other stages develop in visual capillaries. Furthermore, unless several blood surveys are carried out at different time of the year, and are combined with spleen rates, it is not possible to predict with any degree of certainty the amount and intensity of perennial malaria endemicity in a given area (Voller and O'Neill, 1971).

#### 9. Serological method :

Serological tests such as indirect fluorescent antibody (IFA) test and to a lesser extent indirect (passive) hasm-agglutination tests, ensyme linked immunosorbent assay (RLISA), and gel precipitation tests also have a diagnostic role in epidemiological studies. In the past, these tests had suffered disadvantages due to mon-evailability of pure forms of antigens or antibodies.

The recent isolation, however, of purified antigens from all stages of parasite and the development of specific monoclonal antibodies have resulted in the development of more specific reagents and a new generation of tests. These, alongwith the recently identified

parasite-specific DNA probes, have increased the prospects of applying to epidemiological studies new and improved diagnostic tests, capable of automation (W.H.O., 1987).

A great variety of antibodies are produced during the course of malaria infection. These may be detected by precipitation of soluble antigens, fluorescence. agglutination opsomization of parasitized cells, and blocking of merosoite penetration (Playfair, 1978). Opsonizing antibodies (Rogers, 1974), merosoite blocking antibodies (Michell et al. 1975) and delayed hypersensitivity (Phillips et al. 1970) provide protection from reinfection but are slow to develop. The precipitin, fluorescence and agglutinin autibodies rise early during infection and persist for long periods. The role of serological procedure like indirect fluorescent antibody test is very promising in the epidemiologic interpretation of date in malaria (Draper et al. 1972; 1972 by Voller. 1971). The period prevalence of malaria in the community as seen by age related antibody profile has been shown to be a more sensitive tool for surveillance as compared to perasite index which provides point prevalence data (Draper et al, 1972 as Bruce Chawatt et al, 1975; Lobel et al. 1976; Kumer et al. 1986, 1987). IIP provides period prevalence data which was more informative as compared to point prevalence data given by paresite rates (Collins et al, 1967; No Farlane et al, 1970; Neuwissen,

1974; Spencer, 1979). Serologic data seem to be more consistent with parasite indices in stable transmission area (Dreper et al. 1972 b). Kumar et al (1987) reported the use of serological procedures in the measurement of malaris in a community and shown it to be more reliable than conventional methods like the parasite index. The infection rates of malaria in the community as calculated from the serological data confirm the reliability of serology in the measurement of malaria.

Sero-positivity correlates well with parasitanmia more so during the transmission season. The amount of antibodies were higher in the non-transmission season as observed by Rumar et al (1986). Lower antibody titres during the peak transmission could be due to absorption of antibodies by the parasite in the blood at a rate faster than they are produced.

Repeated exposure in endemic areas should result in increase antibody levels which would be reflected in an age related increase (Collins et al, 1967; Draper et al, 1972 b). Therefore, particularly in higher age group, there may not be parasites in blood due to immunity, thus peripheral smear examination has certain limitation.

In view of this, serology seems promising in the areas of high incidence of malarie; the age group above 4 years acquire immunity induced by repeated infections, resulting

more number of unrecentised cases, and therefore, no correlation with overall incidence but within the age groups of 1 - 4 years of age as expected, there was good correlation of antibody status with incidence of malaria. However, since serology cannot distinguish between <u>p.vivax</u> and <u>P. falciparum</u> infections active surveillance by smear examination should continue (number et al., 1987).

The following serological methods have been applied in epidemiology of malaria:

- 1. Engyme linked immuno-sorbent assay (ELISA)
- 2. Indirect Issuno-fluorescence test (IIF)
- 3. Indirect hasmagglutination test (IHA)
- 4. Radio Immunossay (RIA)
- 5. Latex agglutination test
- 6. Gel immunoprecipitation test
- 7. Counter current immuno-electrophoresis (CIEP)
- 8. Complement fixation test.

#### 9.1 Engyme linked immuno-sorbent assay :

performed only in few large centres.

The generic term ensyme immuno-assay are generally known as Ensyme linked immuno-sorbent assay (ELISA). There are two widely accepted assays that employ labelled antibodies and antigens. They are immuno-fluorescence, in which a fluorescent dye is conjugated to the antibody, and radio-immunoassay, in which isotopes are attached to antibodies or antigens. Noth assays are complex and can be

The introduction of ensyme immunoassays, pioneered by Emgwall & Permann (1971) offered an attractive alternative by using enzyme labelled antibody or antigens. The range of application of enzyme immuno assays, is potentially as wide as that of radio-immunoassay (Bull. W.M.O., 1976). The tests though evaluated in different laboratories may not be applicable in the field for diagnosis of malaria at present movement. However, it is envisaged that with the evailability of different specificities of monoclonal antibodies by way of hybridoma technology and also with the help of recombinant DNA techniques immuno-diagnosis of malaria in the field situation may become a reality. Today, precipitation tests and redio-issumo-assays are rarely used, the former because of their sensitivity, the latter, because they have almost completely been replaced by RLISA (W.H.O. Immunodiagnosis in Malaria, Unpublished document, WHO/ Mal. 185, 1018, 1985) .

## Field applications :

For malarie the test was used by Weller et al (1974 a). Since then the test has been used in large number of studies of malaria (Ambroise Thomas et al. 1978; Edrisson at al. 1979; Mahajan et al. 1981; Erivestava, et al 1983, 1981; Ray et al. 1983a, b; Dutta et al. 1982, 1984; Spencer at al. 1979, 1981; Voller et al. 1976 b, 1975.

Marits :- This is a simple technique requiring a limited amount of maternal antique which can be fixed on variety of solid supports from multi-well plastic plates to nitrocellulose paper.

my using purified and defined material antigens in the ELISA, it proved possible to measure in a reproducible way the antibodies against asexual blood stages of <u>F. falciparum</u> in children to increase the sensitivity and specificity and to standardise the method. It can be automated for use in central laboratories where large numbers of samples have to be processed and the results may be quantitated. It may be used under field conditions where the test is semi-quantitative and can be read visually.

Descrite :- The main limitations are inter-laboratory variation due to difficulties in standardization and the relatively poor specificity and sensitivity of the ELISA when perasitized red blood cells extracts are employed for the coating of solid support. The method can be improved by the use of purified antigens (W.H.O. Bull, 65, 1986).

#### 9.2 Indirect Immunofluorescence Antibody (IIV) Test:

This test was introduced by Coons et al (1942).

Since then it has been intensively used in sero-diagnosis of many parasitic and microbial diseases. Brooks et al

(1959) detected P. heghei antibodies by this test and opened a new chapter in epidemiology of malaria.

Washed infected red blood cells used as antiquen. Serum containing antibodies is incubated with antigen. The entigen antibody complex is coupled to a fluorescein labelled antisera and slides are examined by fluorescence microscopy.

The community used antigens have been obtained from patients affected by <u>P. falciparum</u> cultures and monkey blood affected by <u>P. knowlesi</u>, <u>F. cynomolgi</u> and <u>P. costneyi</u>.

#### Pield Application :

(Ambroise Thomas et al. 1972, 1974; Thomas et al. 1975; Carval et al. 1981, 1982; Bruce-Chawatt et al. 1973, 2175; Meuwissen et al. 1974; Collins et el. 1968, 1967, 1971; 1972; Kegan et al. 1981; Voller et el. 1968, 1974; Culzer et al. 1969, 1975; Mehajan et al. 1981; Gupta et al. 1981; Warren et el. 1976; Keay et el. 1973; Mornstein et al. 1983; Ray et al. 1983; Hall et al. 1978).

Srivestave et al (1983) observed its high diagnostic value since 98 percent of slide positive melaris patients carrying P. felciparum or P. vivez could be diagnosed. Furthermore positivity observed in

The preparation of comparable hatches of antigen is relatively simple. The whole infected cell, morphologically identifiable, is used as antigen. The results of the test can be used to show differences in salarie endemicity between localities, and to detect transmission. A higher titres, the test is wirtually always specific for malaria and sometimes can be used to indicate species prevalence. Any laboratory with facilities for carrying out IIF test for other diseases can perform the test for malaria if the antigen is provided.

Malarie parasite carriers can occasionally give negative reactions. This has been especially with children. The necessity for specialized equipment and personel limits, this test to major laboratories. Antigens are available from only a few centres and their storage requires considerable refrigeration space. The transport of antigen can precent problems.

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MATERIAL AND METHODS

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#### MATERIAL AND METHODS

#### 1. The study eres :

The present study was conducted in 20 villages namely - Pahari, Mirona, Maheba, Dhawani, Maral, Bakuan, Babri, Moreta, Chushwan, Gulara, Mod Khurd, Mod Kalan, Sant-Behta, Kibi, Bajehera, Bangra, Simthiri, Jaryai, Chhirona and Sultantpura, located within the area of Primary Health Centre (P.H.C.) Chirgaon which is the rural health training centre (H.H.T.C.) of the Department of Social and Preventive Medicine, Maharani Lammi Bal Medical College, Jhansi (U.P.). The centre is being utilized for field training of undergraduate students in community health and for epidemiological researches.

#### 1-1 Topography :

District Jhansi of Bundelkhand region situated in south-west of Uttar Pradech, is surrounded by districts of Swelior, Datiya, Shivpuri and Tecknoparh of Madhya Pradech and Lalitpur, Hamirpur and Jalaun of Uttar Pradech, P.H.C. Chirgson is situated on Humbay-Eampur road at a distance of 25 kms. from M.L.B. Medical College, Jhansi, U.P. It renders health care delivery to the population of 120 villages besides Chirgson town where centre is located. Hajority of study villages are

connected by pucca roads with the centre; a few, however are not approachable by easy means.

The geographical area of Community Development Block Chirgeon is 55,255 hecture constituting mainly of Padua soil which is suitable for wheat cultivation.

#### 1.2 Climate :

Climate of the area is hot and dry. Mean monthly maximum and minimum temperature ranges between 47.1°C to 3.7°C respectively (1986-1987). General and real rainfall was recorded as 879 mm and 586 mm respectively during the calender year 1986. Mean monthly relative humidity ranges between 15% to 76% at 8730 hrs and 26 to 86% at 1530 hrs (Statistical Diary, U.P., 1987).

#### 1.3 Population composition :

P.H.C. Chirgaon has a population of 1,00,561, according to 1981 census. The density of population is 1.96/hectare. Hele: Female ratio is 1900: 818. The literacy rate is 1:6 higher in comparison to Uttar Predesh and 7.17% lower in comparison to whole of India (Census, 1981). Hajority of them are Hindus followed by Huelima and them others (Covt. of U.P., 1986). Agriculture and labour are main accupations of the area.

#### 1.4 Environmental conditions :

Mostly, houses are either kutcha or semi-pucca with a little or no facility of cross-ventilation. Open and insanitary wells are main source of water supply. There are no sewage and drainage system for disposal of excreta and waste water respectively.

Incidence of malaria was however not uniform throughout the block. These are foci of high and low incidence. The estimated A.P.I. of these villages was over 2.6 per thousand population as reported by District Malaria Office, U.P. (District Malaria Office, Jhansi).

Annual parasite incidence and slide positivity rate in Chirpson block during the year 1982-89 is as follows:

Year	A.P.I.	8.P.R.
1903	12.05	12.27
1983	19.01	12.16
1984	7.32	8.39
1985	7.99	0.16
1986	6.60	0.17
WAY CO.	2.56	2.89
son in the	2.02	2,28
8000	1,04	1.10

Source : District Malazia Office, Jhonsi (V.P.).

FIG. 3.1.

## NO OF SAMPLES COLLECTED FROM EACH VILLAGE (1-20)

	PAHARI MIRONA	เรีเ 91	ON RIVER	<u>.</u> 	TO KANPL	JR.	
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7	GHUSHWAN			M.P.		5 6	
	TAN WAD	-~	<b>)</b>				
	GHUSHWAN	26	$\sim$	15	BAJHERA	65	
	GULARA	46	21	16	BANGRA	80	
	MOD KHURD	34	t: 1855.fel - 121 =	17	SIMTHARI	92	
	MOD KALAN	٧ 54		18	JARYAYI	87	
	SANT BEHTA	28		19	CHHIRONA	156	
	NIBI	38		20	SULTANPURA	498	

FIG. 3.1.

## NO. OF SAMPLES COLLECTED FROM EACH VILLAGE (1-20)

and the second						
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5	BARAL	184		·,	MOTH	1
6	BAKUAN	96		1		
7	BABRI	37	h	1/	. /	
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	SH5.X WAR GHUSHWAN		~~	M.P		7
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69873AS				) 15	BAJHERA	65
10		46		16	BANGRA	80
	MOD KHURD			17		92
12	MOD KALAN					0.00
13	SANT BEHTA	28		19	CHHIRONA	156
14	NIBL	38		20	SULTANPURA	498

#### 2. Study design :

The population survey was carried out once during the transmission period (September and October, 1987).

In this district, there was marked increase in the transmission level during the period September to October as observed by Srivastava at al (1975). The maximum prevalence is from August to November in most of the parts of India.

Seeing the paucity of time and limited resources available, it was thought to conduct the study in transmission period only. The number of samples collected from each village is shown in Figure 3.1.

#### 2.1 Unit of study :

Households of every tenth house selected by systemic random sampling was the unit of study.

#### 2.2 Seepling size and sempling :

In this study every tenth house was the unit of study. All individuals of a household were taken into study issuspective of their ego, sex and health status except for infants under six months of ago, who were not included to svoid the effect of maternal antibodies on the results of sero-epidemiology.

Twenty percent villages in the area under study were selected using simple random sampling method. This was done to provide a 20% sampling of the population of villages under study with inclusion of all age groups. The selection of villages in the block was done by simple random sampling method using table of random numbers (Fisher & Yates, 1957).

The family records of these selected villages as maintained by respective ARN's and CHV's in Chisquon block, varified and made uptodate by making necessary alterations and additions during household listing in the selected villages. The records were re-verified at the time of sampling. The door to door survey was carried out by visiting once during the transmission period (September & October, 1987).

Heads of families of selected household were interviewed on a pre-tested schedule (see Appendix I) to collect information reparding verious bio-social characteristics. Thereafter each individual of household was interviewed separately and information were recorded on a separate schedule (Appendix II). Every individual was examined clinically to find out any associated illness and to assess organomegaly.

2.3 Collection, transportation and Storage of Assolut :

The blood samples of each individual with family

by door to door visit. By finger prick method, two spots of 2 cm. size were taken on Whatman's No. 3 filter paper strips. A thin and thick smear of individual was also prepared. The filter paper strips were air dried in shake. Dried samples were scaled in polythene bags and were transported to the laboratory in ice. In the laboratory, the filter paper strips were stored at -20°C until final analysis. The slides were fixed in methanol on the same day and stained with Geimsa stain. Later, they were examined under oil immersion lens of bisocular compound microscope.

The collection and staining of glass slides were performed in usual manner (W.H.O., 1961). The blood films were stained with Geimsa Stain and examined for malarial parasites.

#### 3. Performance of Serological Test :

#### 3.1 Antigen :

P. folginary entique was proposed from in-vitro sulture of P. folginary entateined at Pational Institute of Communicable Diseases (RICD), Delhi. Test was essentially performed as described by Muli gt gl (1978) and some modifications suggested by May gt gl (1983). The persons was at a sub-culture level of 251 and contained bypromisetely D-7 persons persons with multiply minimum. The autions was prepared by supposing

treatment of the culture followed by sonication. Antigen was schizont antigen and was more than 90% pure.

#### 3.2 Reference sers :

4 3 5 44

The positive reference serum was obtained from a person having heavy malaria infection. The negative reference serum was a pool from slide negative apparently healthy human beings. These had previously been tested by the IIF & ELISA.

- 4. ENZYME LINKED IMMUNOSORBENT ASSAY (ELISA)
- e) The Micro-ELISA test was performed in the 96-well flat bottom polystyrene Micro-ELISA plate (Nunc. Micro-titre No. 1 were used as carrier surface for entiren.
- b) Engywe Conjugate :- Anti-human IgG (heavy and light chains) labelled with horse redish peroxidase was obtained from Cappel Laboratories (Cochran ville, U.S.A.)
- c) Substrate :- Ensyme substrate ortho-physnylene dismine (O.P.D.) was obtained from Sigma Chemical, V.S.A. for measuring paroxidase activity.
- A reference positive and reference negative serum

  were used for determining optimal dilution of antigens,

  merum and conjugate using in-vitro culture <u>l'iniciparum</u>

  antigen dilution from 1 : 3000 and serum dilution

  ranging from 1 : 400 in F.D.S. Tuesn-20.

The optimum antigen dilution using in without culture Po falciparum antigen that gave a strong reading with positive serum and low reading with similar dilution of negative serum was found 1: 3000 and serum dilution was 1: 400. The optimal conjugate dilution was determined by chaquer board titration. Conjugate dilution (1: 1500) was found to be optimum.

Micro-ELIBA Procedure :- Test was performed according to method described by Ray et al (1983).

#### (i) Sensitization of Micro-titre plate :

Antigen was diluted to optimum concentration in coating carbonate buffer (0.06 M, PH 9.6). 200 ul each of the optimally diluted antigen was added into the wells of a micro-titre plate. The plates were covered and hept in plantic box to incubate at 4°C for 18 hrs.

Dilutions (1 : 200 ul) of the test and control sers were made in P.B.S. pH 7.2 containing 0.05% Tween-20 (see Appendix III). The antigen sensitized plates were task out from the refrigerator and the excess liquid was removed from the plate. The plate was washed twice in seme buffer for 5 minutes each time and then dried.

#### (11) Insubstion with sern :

The entigen coated wells were filled with test and control sera of 200 ml on the washed antigen sensitized plate. Antigen control was kept adding P.B.S. Tuesn-20 only. The plates were covered and incubated at room temperature incide the wet plastic box for one hour. The wells were washed for S minutes thrice with P.B.S./T to remove unbound serum.

- (iii) Incubation with Conjugate: Each of the well was then filled with 200 ul vol. of optimal diluted (1: 1500) conjugate and then incubated in a humid chamber at room temperature for one hour. After incubation unbound conjugate was removed with washing three times P.B.S./T
- (iv) Substrate reaction: Peroxidese bound to the wells was finally estimated by adding 200 ul of substrate solution in each well and incubating the plate at room temperature in the dark. The reaction is allowed to proceed for 10-15 minutes. The reaction was stopped with the addition of 50 ul of 58 H<sub>2</sub>SO<sub>4</sub> in each well.

For expression of results, the reading at a dilution of 1 : 400 was used since at this dilution, the positive-negative differentiation was best. 93-4 percent of the control sera from Delhi and all the sers from Eachmif gave a negative ( $\angle$  0-4) reading. Taking this as the point of differentiation between the positive and negative sers. \$2.50 percent of our study individuals showed reaction upto  $\theta = 0.6$  0.D. (2492); 24.01 percent showed 0.4 = 0.6 0.D. reaction (2492); 14.01 percent

individual showed 0.6 - 0.8 O.D.  $(E_{492})$  and 5.05 percent showed 0.1 - 1.0 O.D.  $(E_{492})$  reaction. All the slide positive individual showed more than 1.0 O.D.  $(E_{492})$  reaction.

Reading was taken at optical density (00) at wave length of 492 n.m. using spectronic 20 spectrophotometer

#### 5. Indirect Immunofluorescence Test :

A local strain of 2-falciparum (FAN-5) was adopted to continuous culture and maintained in NICD Leboretory since 1978 by the method of Rai Chowdhuri et al.

At sub-culture level of 122 when the parasitaenia was 8% with rings (20 percent), trophosoites (35 percent) and schizont (45%), the culture was washed five times in P.B.S.

pH 7-2. After the final washing, it was suspended in the same buffer in such a way so as to contain about 20-30 plasmodia per high power field in a thick swear (Sulmer et al., 1969). While preparing the swears, care was taken that the cells did not settle out of the antigen suspension in the pasteur pipettes. After drying, the smears were stored at -70°C, wrapped in was papers.

#### S-1 Reference sere :

The reference malaria positive sera from salaria cases and negative sera from non-malarious area were secrived from C.D.C. Atlanta and stored as above.

#### 5.2 Fluorescent Conjugate :

Anti-human IgG (heavy and light chains) labelled with fluorescein isothiocynate was obtained from immuno-diagnostic limited. Different conjugate dilutions were tested for finding the optimum dilution to be used. It was found that conjugate dilution 1 : 10 was giving highest titre with reference positive and lowest with reference negative sers.

#### 5.3 Performance of the Test :

method of Sulzer <u>et al</u> (1969) with some modifications suggested by Ray <u>et al</u> (1982). The antigen slides (stored at -70°C) were taken out and kept on racks made in glass petridishes and were labelled and allowed to dry. The test sera along with positive and one negative control sera were diluted in two fold dilution starting from 1:32 to 64 in P.B.S. pH 7.2. A drop of each dilution of test sera was placed covering each antigen smear. A control smear was label receiving P.B.S. pH 7.2 instead of serum. The slides were placed inside humid petri dishes and incubated at 70°C for 30 minutes. Next the slides were washed thrice (each time for 10 minutes with P.B.S. pH 7.2) with manual stirring and dried quickly under the fams.

Optimal dilutions (1 : 40) of commercial antihuman IgG, A and M (N and L) conjugated with fluorescin isothiocynate (Institute Pasteur Production) was added to cover the smear fully.

Incubation, washing and drying in the above manner followed. The slides were mounted with buffered glycerol (pH 7.2) and examined under a fluorescent microscope.

#### 5.4 Reading and interpretation of Results :

Pluorescence was subjectively graded from negative to 4+ and ++ and above were considered positive.

The fluorescence of the parasites were seen against a background of faintly visible erythrocytes (Rey et al. 1982).

#### 6. Compilation, Tabulation and Interpretation of Date :

Data so obtained from the study was subjected to exitical statistical analysis which consist of estimation of the prevalence of antibody titre in random population and to find out, correlating it with various bio-social characteristics of the population.

The total tests of significance such as Chi square took was used to determine the significance of the association between the two variables and difference between two parametric values.

#### 7. Limitation of study :

The study had been carried out in partial fulfilment of the requirements of N.D. (Social & Preventive Medicine) examination and therefore suffers from limitations of time and resources. Many of the informations sought, are based on the capacity to recell, the limitations of which do not need any emphasis. The reluctance on the part of individuals in giving the blood samples proved a great difficulty in the course of study. Inspite of the best efforts made, such samples of all individuals could not be obtained.

To show seasonal variations in the transmission of the disease, the non-transmission survey could not be conducted due to paucity of time and resources available.

Due to unavoidable reasons and paucity of time. the IIP test could be performed only in ELISA positive proven samples of blood.

#### 8. <u>Different criteria adopted</u> :

#### 0.1 Family type :

Any family with husband, wife and their offsprings was considered as nuclear and rest were considered as joint.

#### 8.2 Family Size :

A family upto 5 members was considered as small, whereas one with 6 or more members was taken as large.

### 8.3 <u>Social class</u> :

Social classification of families used in this study was as given by Szivastava et al (1982). Criterion of social classification broughtforth by Szivastava et al (1982) is given below :

	an monthly per capite income	Social Class
<b>1</b> •	600/- and above	2
<b>8.</b>	300/- to b. 599/-	11
	140/- to m. 299/-	111
<b>10.</b>	60/- to m. 139/-	17
4	B. 60/-	

\*\*\*\*\*

# 

#### OBS ERVATIONS

The present study was conducted at Primary Nealth Centre, Chisgaon, Jhanei (U.P.). The study was carried out in twenty villages. The systemic sampled population under study consisted of 1520 individuals of 269 families out of total 290 families.

#### 1. Population under study :

There were 98.9% Hindus and 1.09 percent Muslims. Out of these, 15.53% individuals belonged to upper ceste and 49.20% backwards and 25.27 percent scheduled caste. The majority of families were joint (75.29%) and rest belonged to nuclear (24.71%) families. The maximum percentage (32.71%) of families consisted of 5-6 members and minimum percentage (1.42%) of families consisted of 1-2 members. There were three-fourth families in social class IV (46.29%) and class V (28.25%) and one-fourth families classified in social class III (22.30%) and social class II (4.47%). The main occupation of the families was agriculture (71.92%) followed by labour (25.46%), service (2.0%) and rest were engaged in business and other occupations. In the study, married individuals were 59.74 percent and 37.56 percent unmarried and rest of them were widow/widower and divorces. There were

64.54 percent illiterate individuals, followed by literate (19.81%), just literate (5.98%) and children (9.67%).

#### 1.1 Male Pemale ratio :

The mex variation in the study revealed that adult male contributed 36.91 percent and adult female 31.37 percent. The paediatric population 0 - 14 years was 31.72 percent; male accounted for 18.03 percent and female 13.69 percent.

An attempt has been made in this study to see
the relationship between the prevalence of malazia and
its various bio-social characteristics. The elaborate
description for this relationship has been given in
subsequent text, taking the various variables one by one.
The impact of these variables on the distribution of
disease has been viewed separately amongst the studied
population.

TABLE 1 Distribution of individuals by their age and sex.

Age (year)		No. X		70.		Total X	
_	1	1	0.07	**	***	1	0.07
1	- 6	63	4.14	41	2.70	104	6.94
5	- 9	103	6.78	95	6.25	198	13.03
10	- 14	107	7.04	72	4.74	179	11.70
15	- 24	174	11.45	147	9.67	321	21-12
25	- 34	134	8-82	112	7.37	246	16.19
35	- 44	96	6.32	81	5.33	177	11.65
45	- 54	70	4.60	80	5.26	150	9.06
55	- 64	61	4.01	44	2.89	103	6.90
65	•	26	1.71	23	0.86	39	2.57
Tot	<b>al</b>	635	54.93	683	45.07	1520	100.00

Table 1 shows the age and sex distribution of study population.

The percentage of males and females was 54.93 and 45.07 respectively. It was observed that maximum (21.12%) individuals helosped to the age group 15 - 24 years. followed by 16.19 percent in age group 25 - 34 years. whereas, there was only one (0.07%) individual in age

group \_ 1 year. The passistric population encounted for 31.72 percent.

#### 2. Bio-social characteristics of population :

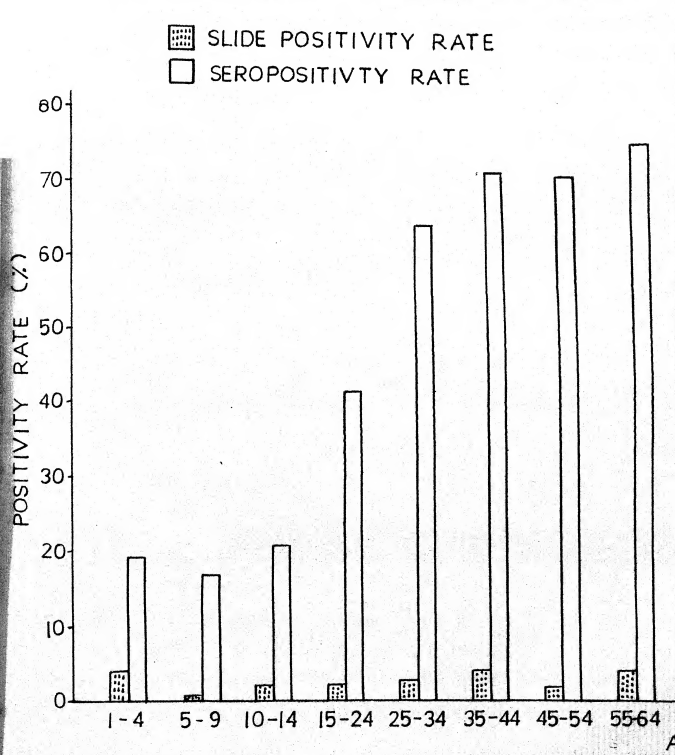
#### 2-1 Aug :-

TABLE 2
Slide positivity and sere-positivity according to age.

(years)		Total exemined	Slide examination No.found Positi- positive wity positive rate(%)		Ferelogical Po. found positive	Positivity Pote (%)	
4	1		•		•	•	
1 -	4	106	•	3.05	20	19.23	
5 -	9	190		0.30	33	16.66	
10 -	34	179		1.60	37	20.67	
15 -	24	321	6	1.07	134	41.74	
25 -	34	246	11	4.47	150	64.22	
35 -	44	177	7	3.95	126	71.19	
45 -	54	189		2.00	100	70,66	
55 -	64	105	•	3.01	79	75.24	
65 +		39	•	2,56	•	74.36	
retel				2,63	123	47.50	

(2 -2.77, a.e.-2. > C.0.25) (2 -300.1, a.e.- a. > C.0.001)

FIG.- I
BAR DIAGRAM SHOWING S.P. R. SEROPOSITIVIT
RATES AMONGST INDIVIDUALS BY AGE



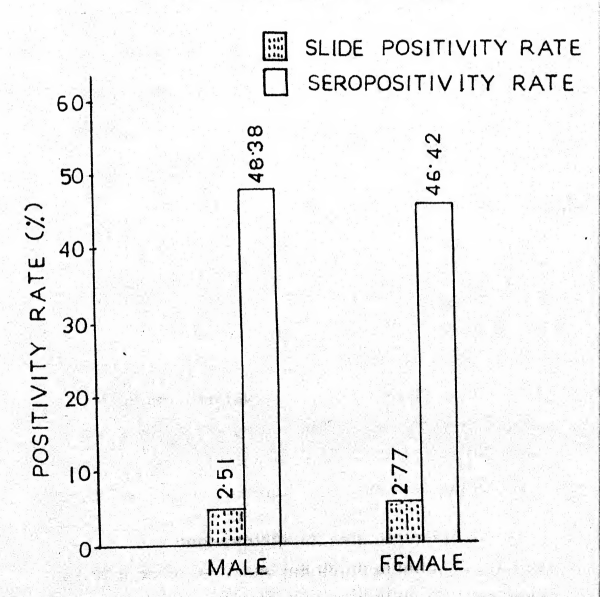
AGE GROUP (YEARS)

Table 2 and Fig. 1 shows that out of total 1520 individuals, 40 (2.63%) were slide positive for malaris. P. vivas infection was detected in all these cases. The slide positivity rate was highest (4.47%) in sqe group 25 - 34 years, followed by 3.95 percent in the age group 35 - 44 years. The higher (3.85%) positivity in age group 1 - 4 years shows that fresh transmission is occurring in this area, difference is not statistically significent.

The sero-positivity rate of 19.23 percent
was seen in \$\left( 1 - 4 \) years age group, 20.67 percent
in 6 - 15 years age-group, 71.19 percent in 35 - 44
years age-group and 75.24 percent in 55 - 64 years and
above age groups.

those aged (\_ 1 - 14 years when compared with 15 - 44 years and 45 - 64 years age-groups.

FIG.-2
BAR DIAGRAM SHOWING SLIDE POSITIVITY
AND SEROPOSITIVITY RATES AMONGST
INDIVIDUALS BY SEX



Constitution of the second

TABLE 3
Distribution of slide positivity and sero-positivity according to sex.

	Total No. examined	Slide e No. found positiv	Positivity rate (%)	Serologica No. found positive	Positivity rate (%)
Mal•	835	21	2.51	404	48.38
Pemale	<b>683</b>	19	2.77	319	46.42
Potal	1520	40	2.65	722	45.50

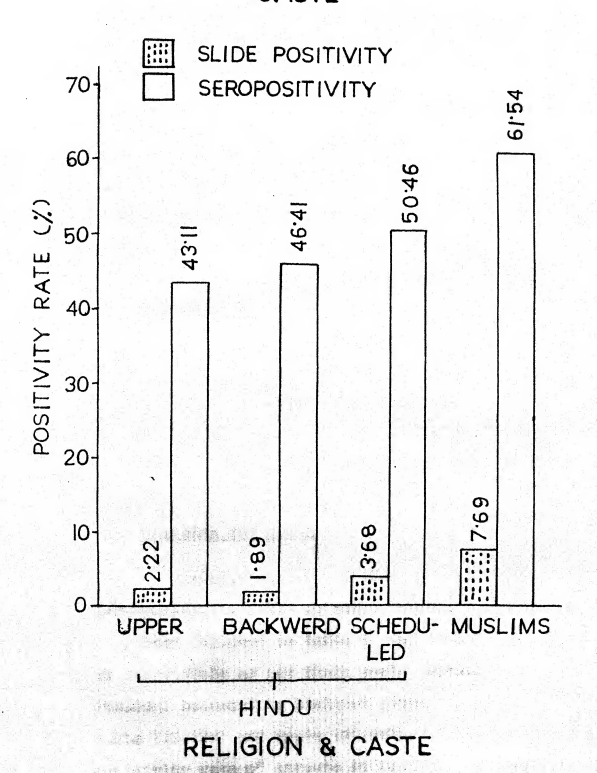
(x2-0.1, e.f.- 1. » (0.75), (x2-0.583, e.f.-1, » (0.25)

### 2.2 845 .

Table 3 and Fig. 3 shows the distribution of the individuals according to their sex. The percentage of males and females is 54.03 and 45.07 respectively (male : female :: 1000 : 618). The slide positivity rate was 2.51 percent in makes and 2.77 percent in females. The difference is not statistically significant.

The seco-positivity rate was observed as 40.38 percent in makes and 44.42 percent in families respectively. He significant statical difference was observed in both sexus.

FIG. - 3
BAR DIAGRAM SHOWING SLIDE POSITIVITY &
SEROPOSITIVITY ACCORDING TO RELIGION AND
CASTE



Slide positivity and sere-positivity according to religion and caste.

Religion	No.of		Slide examination		cal ion
4 easte	Cases	No. found positive	Positi- vity	No.	Positi- vity rate(%)
Beligien :					
. Mindu :					
- Upper	225	5	2.22	97	43.11
- Rackward	739	14	1.09	343	46.41
- Scheduled	543	20	3.68	274	50.46
. Muslims	13		7.69	•	61.54
Total	1820	40	2.43	722	47.50
(x <sup>2</sup> =4.046, d.f.	2. 2	79-10).	(x <sup>2</sup> -4.000.	d.f.=3, P	70.25

#### 2.3 Religion and coste :

Distribution of individuals, slide positivity and seco-positivity rates by their various soligious and castes, have been deploted in table 1 and fig. 3. The distribution of individuals as per their caste remealed that maximum (48,699) belonged to beckward caste, followed by scheduled easts (35.729) and upper (16,600). While calculating the positivity rate of disease is relation to costs, it was

Service of and the first the service of the service

observed that all individuals were Hindu except one slide positive case being Huslim.

However, the slide positivity rate was highest (7.69%) for Muelims, whereas for scheduled and backwards it was found to be 3.68 and 1.89 percent respectively.

The difference observed was, statistically not significant. The sero-positivity rate was also higher in Muslims (61.54%) and scheduled castes (50.46%) followed by backwards (46.41%), it was lower (43.11%) in upper castes. However, difference was, statistically insignificant.

Slide positivity and sero-positivity according to their merital status.

Meritel Status	Total		ide netion	Serela	gigal etion
		So. found possibility	Foalti-	No. found positive	Positi-
Natried	200		3.19	565	62.22
Venerried	571	11	1.99	130	22.77
Widow/ Widowez/ Divozceo			<b>\</b> . <b>\</b>	•	65.85
Sotal .	850	•	2.40	723	47.50

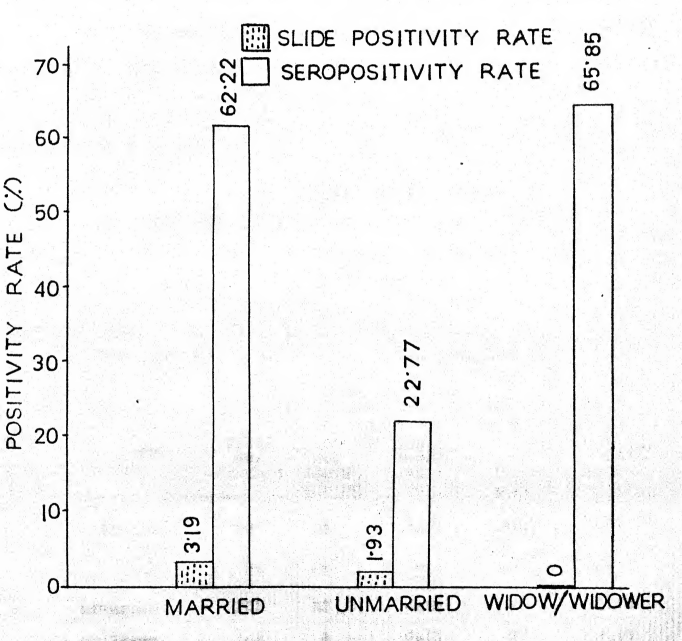
(x2-8-95, d.f.-1, p (0.01), (x2-224-4, d.f.-2, p (0.001)

FIG. -4

BAR DIAGRAM SHOWING SLIDE POSITIVITY

AND SEROPOSITIVITY AMONGST INDIVIDUAL

BY MARITAL STATUS



MARITAL STATUS DIVORCE

and the first to the

ESTABLISHED SHADO SE DERKIN

# 2.4 Marital Status :

and positivity rates of malaria in relation to their marital status. The majority (59.74%) of individuals were married, followed by unmarried (37.56%). The widow/widower and divorces were 2.69 percent. The slide positivity rate was higher (3.19%), in married and lower (1.93%) in unmarried individuals. The difference was statistically significent. The sero-positivity rate was higher (63.65%) in widow/widower and divorces and in married (62.22%) and it was lower (22.77%) in unmarried individuals. The difference was statistically significent.

Slide positivity and sere-positivity according to literacy status.

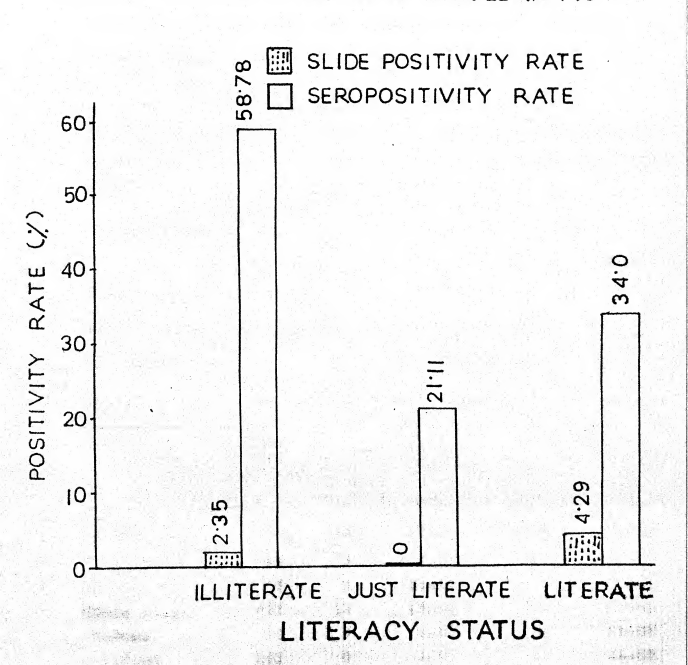
	Total	Slide exemination		Serological exemination	
Literacy status	examined	No. found positive	Positi- wity mate (%)	found positive	Post ti- vity Fate (%)
Illiterate	•••	29	2,35	576	50.70
Suet Literate	•	•			21-11
Messe			4.39	103	34.00
children	147	ė.	2.73	*	16.33
	1525		2.00	723	47.50

(x2-4-126, 4-1-1, p 70-025), (x2-191-601, 4-1-2, p\_0-001)

FIG. -5

# BAR DIAGRAM SHOWING SLIDE POSITIVITY RATE & SEROPOSITIVITY RATE AMONGST INDIVIDUALS BY LITERACY STATUS

( CHILDREN UPTO 5 YRS, WERE IGNORED IN FIG )



# 2.5 Literacy status :

Relationship between literacy status of individuals and slide positivity and sero-positivity have been shown in table 6 & fig. 5. The majority of individuals were illiterate (64.54%) or just literate (5.98%). Literate contributed only 10.00 percent.

The slide positivity rate of 4.29 percent was observed in literate, followed by children (2.72%). It was lower (2.35%) in illiterate. The difference was statistically significant.

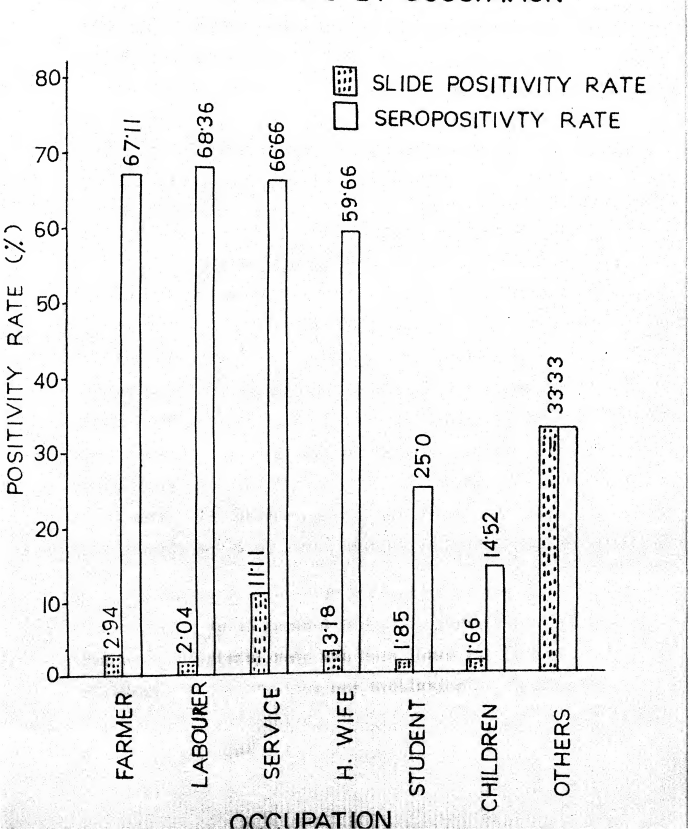
The sero-positivity rate was highest (50.78%) in illiterate and lower (34.00%) in literate. It was further observed that sero-positivity rate declined with improvement in literacy status. However, it was statistically significant.

TABLE 7
Slide positivity and sero-positivity according to eccupation.

Scrupation		Slide examination		Serological exemination	
		No. found positive	Fostel- vity rate(%)	No. found positive	Poelti- vity Este(%)
Farmer	374	11	2.94	251	67.11
Labourer	90		2.06	67	60.36
Service			11-11	•	66.66
Nomes vife	474	15	13-10	201	59.66
Student	326	•	1.65	81	25.00
Children	201		1.44	38	14.52
Others			33,33		33.39
TOTAL		4	2.43		47,50

(x<sup>2</sup>-1-4172, 4-2-2, 2<u>(0.5)</u>, (x<sup>2</sup>-243-927, 4-2-5, 2<u>(0.001)</u>

FIG.-6
BAR DIAGRAM SHOWING S.P. R. & SEROPOSITIVITY
INDIVIDUALS BY OCCUPATION



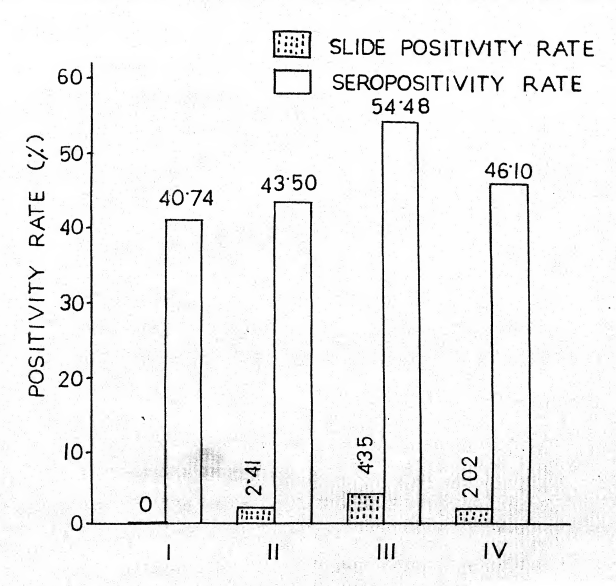
#### 2.6 Occupation :

rate and sero-positivity according to occupation. Farming (24.61%) and labourer (6.45%) were the predominant occupation is the study, followed by housewives who contributed 30.99 percent. There were 21.32 percent individuals amongst students whereas 15.86 percent accounted for children.

The slide positivity rate was higher (11-11%) individuals engaged in service. Nost of them were employed in Parichha Thermal Power Project, while studying population for clide positivity as per their occupation it was recorded that it was higher (2.94%) smonget farmer's and/or housewives (3.18%), followed by 2.04 percent in lebourer. Children showed lower clide positivity rate. However, a very high (33.33%) slide positivity rate was observed in individuals grouped as others. The difference between slide positivity rate for various occupation was statistically not significant.

The sero-positivity rate in labourers and farmers were 60.36 percent and 67.11 percent respectively. The sero-positivity rate was much lower (16.52%) in abildren. The difference was statistically significant.

FIG.-7
BAR DIAGRAM SHOWING S.P. R. & SEROPOSITIVITY
RATE AMONGST INDIDIVIDUALS BY SOCIALCLASS



SOCIALCLASS

TABLE 8

Slide positivity and sero-positivity according to social class.

Social	Total No.	#Ride		Serological examination	
cles	examined	found positive	Positi- vity Fate (%)	No. found positive	Positi- vity
	- 54	•	-	22	40.74
III	331	•	2-41	144	43.50
ΧΨ	391	17	4.35	213	54.40
•	744	15	2.03	343	46.10
Total	1520	40	2,63	722	47,50

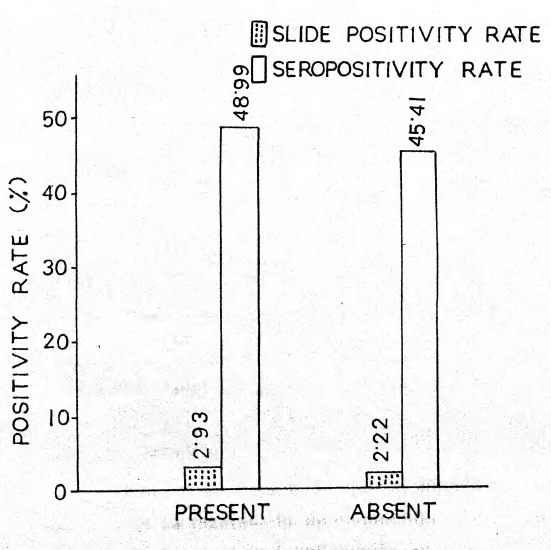
Note: There was no family from Social Class I.  $(x^2-7.125, 4.6.-3, 7.70.05), (x^2-11.262, 4.6.-3, 7.70.010)$ 

### 2.7 Social Class :

Table 8 and 71g. 7 give distribution of individuals in various social classes of population. Out of 1520 individuals, maximum (40.95%) were belonging to social class IV, followed by social class V (15.72%), social class IX (21.78%) and social class XX (3.85%). He individual belonged to social class I. While calculating slide positivity rate was higher in social class V (4.35%) in comparison to social class IV

FIG.-8

BAR DIAGRAM SHOWING S. P. R. & SEROPOSITIVITY
RATES AMONGST INDIVIDUALS BY OVERCROWDING



OVERCROWDING

(2.02%) and social class III (2.41%). Yet the difference was statistically not significant. The sero-positivity rate was also higher in social class v (54.48%) followed by social class IV (46.10%) and lower in social class II (40.74%). However, the difference was statistically significant.

TABLE 9
Slide positivity and sero-positivity according to overcrowding.

	Total	811		Serological exemination		
orouding	No. exemined	No. found positive	Positi- vity	No. found positive	Positi- vity rete(3)	
Present	993	26	2.93	435	40.99	
Absent	632		2.22	307	45,41	
otal (	1520	40	2.63	722	47.50	

# 2.0 Own F-street and 1

Table 9 and Pig. 8 is showing distribution of individuals is relation to over-crowding. Over-crowding was seen in 50.42 percent individuals, who rees (1.56) percent individuals were residing in sufficient number of living manner. The slide positivity rate was almost equal in both groups whether over-exceeding was present or mat.

Slide positivity rate for individuals living undercrowded and un-crowded conditions were 2.93 percent and 2.22 percent respectively; the difference being statistically insignificant.

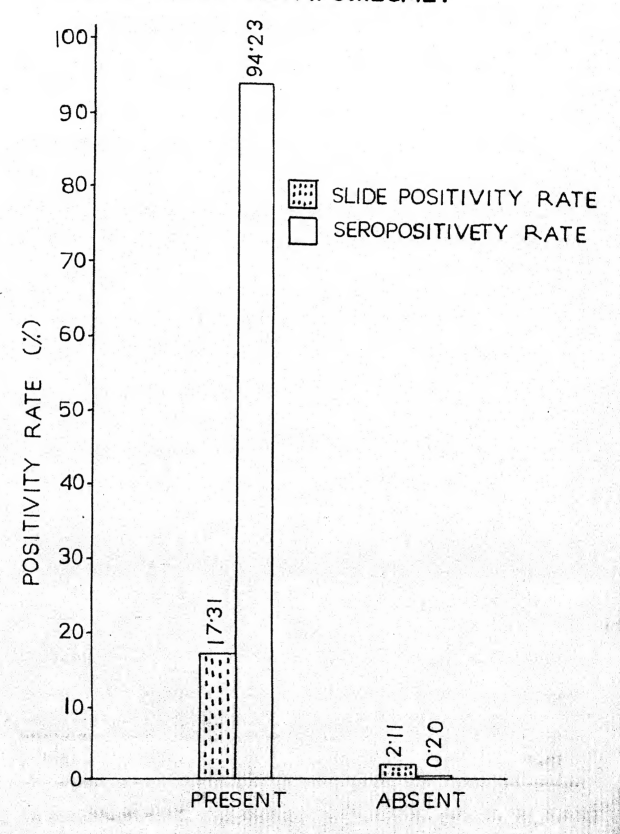
The sero-positivity rate for individuals living in crowded and uncrowded dwelling were 48.99 and 45.41 percent respectively. The difference being statistically insignificent.

TABLE 10 Slide positivity and sero-positivity according to hopetomogaly.

Hepato-	Total	Slide exemination		Seroleg extenine	
megaly	Mo. examined	No. found	Positi- wity rate(%)	No. found positive	Positi- vity
Pzesent	52	•	17.31	49	94.23
Absent	1460		2-11	•	0.20

(x2-03.93, d.f.-1, P (0.001), (x2-1320.02, d.f.-1, P(0.001)

FIG. - 9
BAR DIAGRAM SHOWING S.P.R. & SEROPOSITIVITY
RATE AMONGST INDIVIDUALS WITH HEPATOMEGALY
AND WITHOUT HEPATOMEGALY



HEPATOMEGALY

3. Slide positivity and sero-positivity in relation with clinical manifestations :

### 3.1 Mepstomegaly :

Table 10 and Figure 9 shows, that out of 1520 individuals, 3.42 percent showed hepatomegaly and slide positivity rate was also higher (17.31%) in hepatomegalic individuals. It was lower (2.11%) in non-hepatomegalic individuals. The difference was statistically significant.

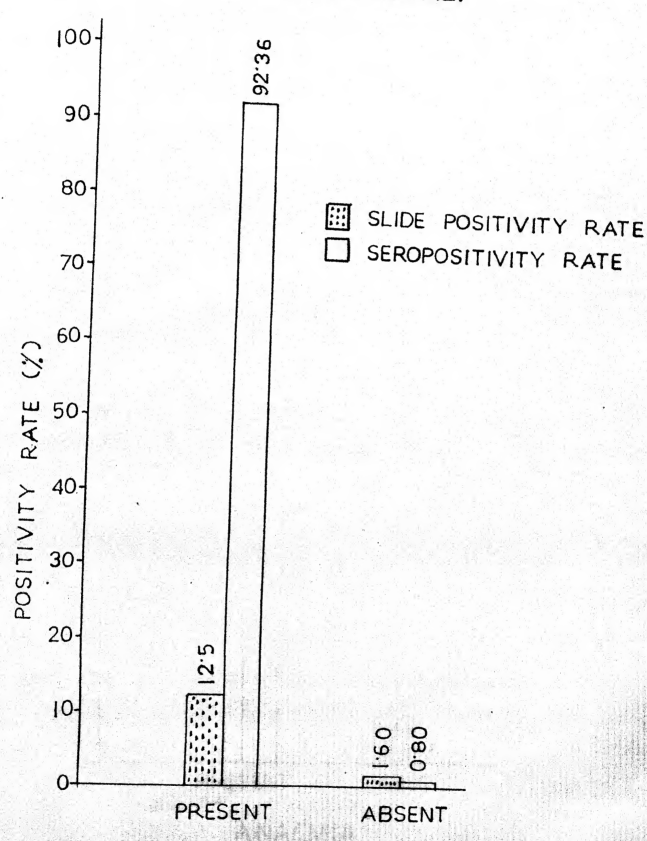
The sero-positivity rate was highest (94.23%) in individuals with hepatomegaly and lower (0.20%) in individuals without hepatomegaly. The difference was statistically significant.

TABLE 11 Slide positivity and mero-positivity according to Splenomegaly.

Degaly committed found vity found positive Estels) positive for found positive for a found		Total			Serological exemination	
	27.47.	80*	Sound	Positi- vity	No. found	Positi-
Ament 1176 22 1.60 11		144	•	12.50	133	92.36
		<b>W</b>	88	1,60		0.00
			•	9.00	144	947

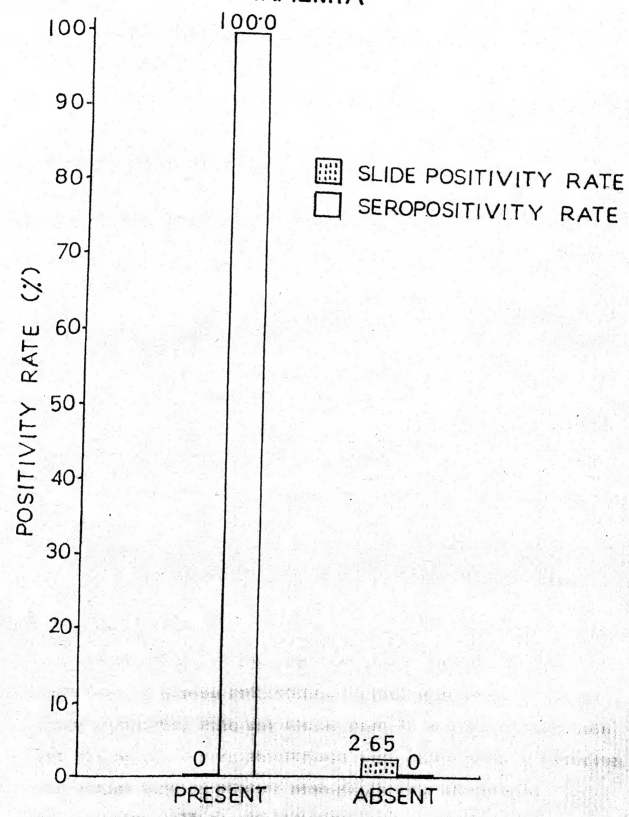
(x2-co.22, d.f.-1, p \_0.001). (x2-1270.3, d.f.-1, p\_0.001)

BAR DIAGRAM SHOWING S.P. R. & SEROPOSITIVITY
RETE AMONGST INDIVIDUALS WITH AND WITHOUT
SPLENOMEGALY



SPLENOMEGALY

FIG.-11
BAR DIAGRAM SHOWING S.P.R. & SEROPOSITIVITY
RATES AMONGST INDIVIDUALS WITH & WITHOUT
ANAEMIA



ANAEMIA

# 3.2 Splenomegaly :

Table 11 and Fig. 10 shows that out of 1520 individuals, 9.47 percent presented with splenomegaly slide positivity rate was higher (12.50%) among individuals with splenomegaly and lower (1.60%) in individuels without splenomegaly. This difference was statistically significant.

TABLE 12 Slide positivity and sero-positivity according to anaemia.

	Total	3 L		Sero logical	
Ansenia	exemined	No. found positive	Positi- vity rate(%)	No. found positive	Positi- vity rate(%)
Fresent				11	100.00
Absort	1509	40	2.45	711	
Total	1520	•	2.63	723	47.50

# 3.3 Appenia :

Individuals, slaves individuals (0.72%) were found amounts.

Filds positivity rate was almost zero in ensemble individuals.

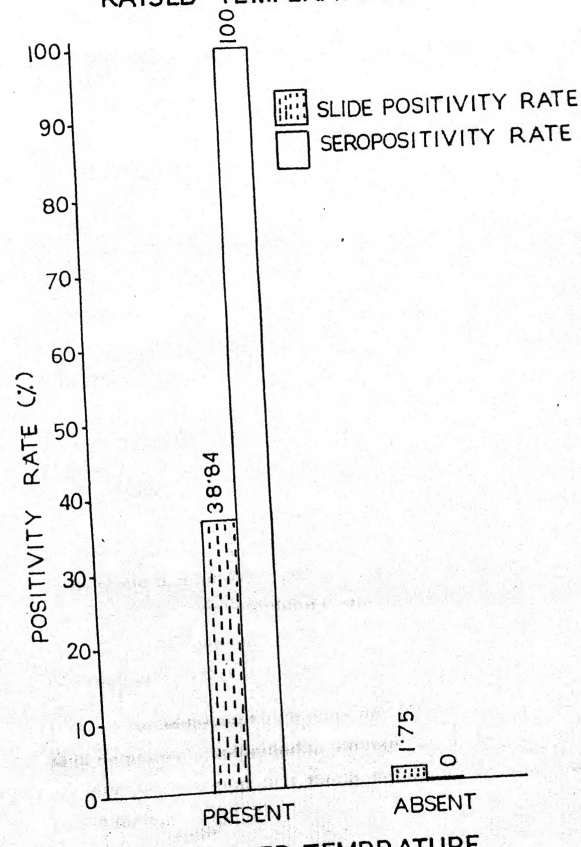
Put all of the cloves individuals were corologically positive.

and should handred pareset sero-positivity is amounts.

Individuals. All the 46 individuals with slide proves.

Parallegais 316 hat them succels at the time of correspond and all all positivity is amounted individuals.

FIG.- 12
BAR DIAGRAM SHOWING S.P.R. & SEROPOSITIVITY
RATE AMONGST INDIVIDUALS WITH & WITHOUT
RAISED TEMPERATURE



RAISED TEMPRATURE

TABLE 13 Slide positivity and sero-positivity according to high

Total	<b>心</b> 发表的工	netion	Serological examination	
No. exemined	found	Positi- vity	No. found positive	Positi- vity
30	14	36.94	30	190.50
1402	26	1.75	684	
1520	10	2.63	722	47.50
	94 1482	examined found positive  36 14 1482 25	examined found vity positive rate(%)  38 14 36.84  1482 26 1.75	examined found vity found positive rate(%) positive 28 14 36.84 38 1482 26 1.75 686

 $<sup>(</sup>x^2-173.567, d.f.=1, P <math>\angle 0.001)$ .

### 3.4 Raised Temperature :

temperature.

Puble 13 and Fig. 12 shows that the slide

positivity and sero-positivity according to high temperature.

In small number (2.5%) of individuals the history of high

temperature was recorded during survey. The slide

positivity rate in individuals with high temperature was

higher (36.86%) whereas it was lover (1.75%) in afabrile

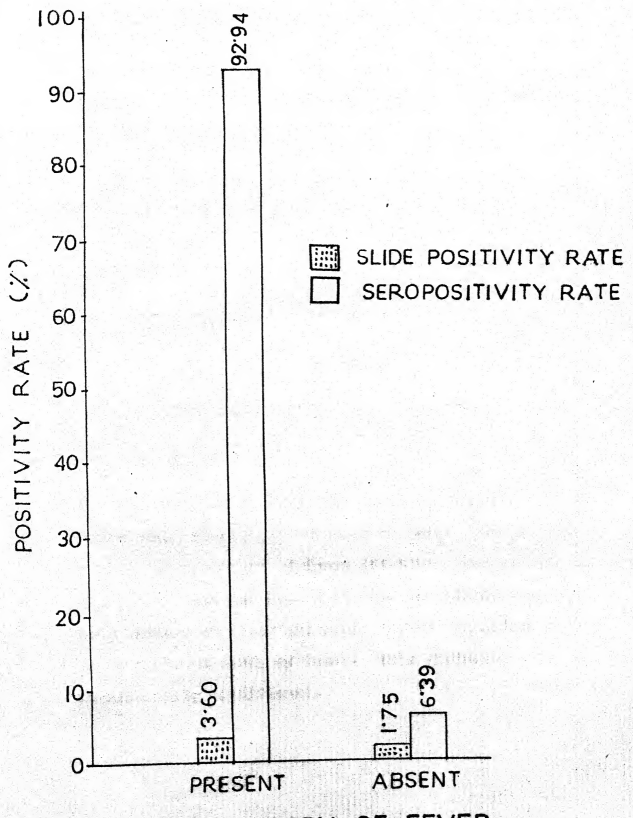
individuals. This was statistically significant.

The sero-positivity rate was hundred percent in block temperature individuals, whereas high temperature was not observed oven in a single secologically negative individuals.

HART HISTORY OF FEMALE

FIG.-13

# BAR DIAGRAM SHOWING S.P.R. AND SEROPOSITIVITY RATES AMONGST INDIVIDUELS WITH AND WITHOUT PAST HISTORY OF FEVER



PAST HISTORY OF FEVER

TABLE 14 Slide positivity and sero-positivity according to past history of fever.

Past Total		00 00 000 000	Slide exemination		gical ation
history of fever	No.	No. found positive	Positi- vity	No. found positive	Positi- vity
Present	722	26	3.60	671	92.94
Absent	698	14	1.75	51	6,39
Potal	1520	40	2.63	722	47.50

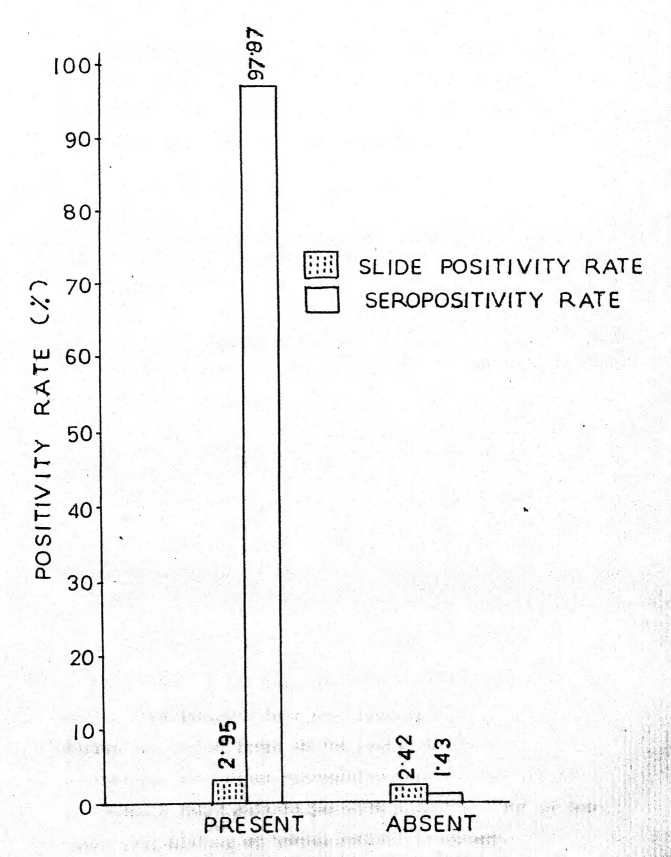
(x2=5.065, d.f.=1, p \( 0.10), (x2=1330.06, d.f.=1, p\( 0.001)

#### 4. Past Mistory :

# 4.1 Past history of fever :

Table 16 and 71g. 13 shows the distribution of individuals having past history of Sever (47.50%) and without past history of Sever (52.50%). The slide positivity rate was higher (3.60%) in individuals with past history of Sever and lower (1.75%) in individuals without past history of Sever. This difference was statistically significant.

BAR DIAGRAM SHOWING S.P. R. AND SEROPOSITIVITY RATE AMONGST INDIVIDUALS WITH & WITHOUT PAST HISTORY OF TREATMENT



PAST HISTORY OF TREATMENT

The sero-positivity rate was higher (92.96%) in individuals suffering from fever at the time of survey and have suffered in past and it was lower (6.16%) in individuals without past and present history of fever. This difference was statistically significant.

TABLE 15 Slide positivity and sero-positivity according to past history of treatment taken (presumptive/redical).

Past Mistory	No. of	SILE			egical maties
of treatment Precumptive/ radical	exemined	No. found positive	Positi- vity rate(%)	No. found positive	Poslal- vity rate(S
Treetment taken	611	10	2.95	598	97.67
Treatment not takem	909	22	2.42	124	1.43
Total	1520	40	2.63	722	47.50
•		. 961 /2		A.f.el.	p/ 0.001

(x =0.385, d.2.=1, P /0.75).

# 4.2 Past history of treatment (Presumptive/redical):

Table 15 and Fig. 14 shows that the majority of individuals (40.20%) have past history of treetment taken during attacks of fever in the past. In this all the individuals have taken presumptive treatment except few individuals found malaria parasite positive. But no one could give history of taking redical treatment.

The slide positivity rate was higher (2.95%) in individuals with past history of treatment taken (presumptive) and lower (2.42%) in individuals without past history of treatment taken. This difference was statistically not significant.

The sero-positivity rate was higher (97.67%) in individuals with past history of treatment taken and lower (1.43%) in individuals without past history of treatment taken. This difference was statistically eignificant.

# Correlation between slide positivity and sero-positivity :

Out of 1520 individuals examined, the forty individuals (2.63%) were detected positive for malaria parasite. Plasmodium vivas infection was observed in all positive individuals. During the same time, the sero-positivity was observed as 47.5 percent. In the forty individuals, all were also found positive sero-logically. Thus shows hundred percent sero-positivity in slide positive individuals. The sero-positive rate positively correlated with the slide positivity rate (Fig. 15).

FIG.- 15
CORRELATION BETWEEN SLIDE & SEROPOSITIVITY
RATES

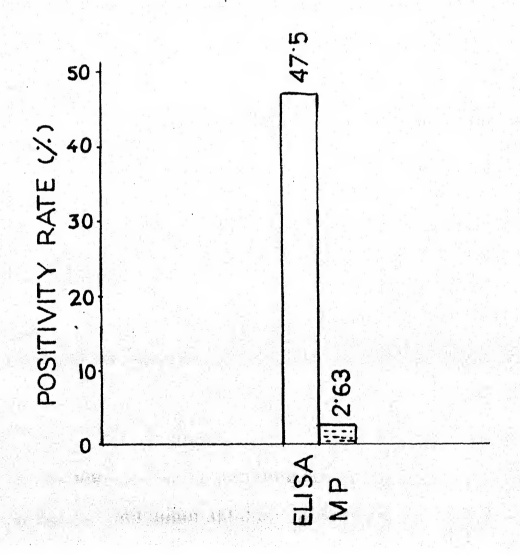


TABLE 16

Distribution of children (2 - 10 years) according to their sex.

5.62	Total No.	Slide examination (2-10 years)		Serological examination (2-10 years)	
	exemined	No.found positive	*	No.found	3
Nalo	189	3	1.59	25	13.23
Penale	139	3	2.16	23	15.40
Total	339	6	1.03	67	14.33

### 5. Paresite rate :

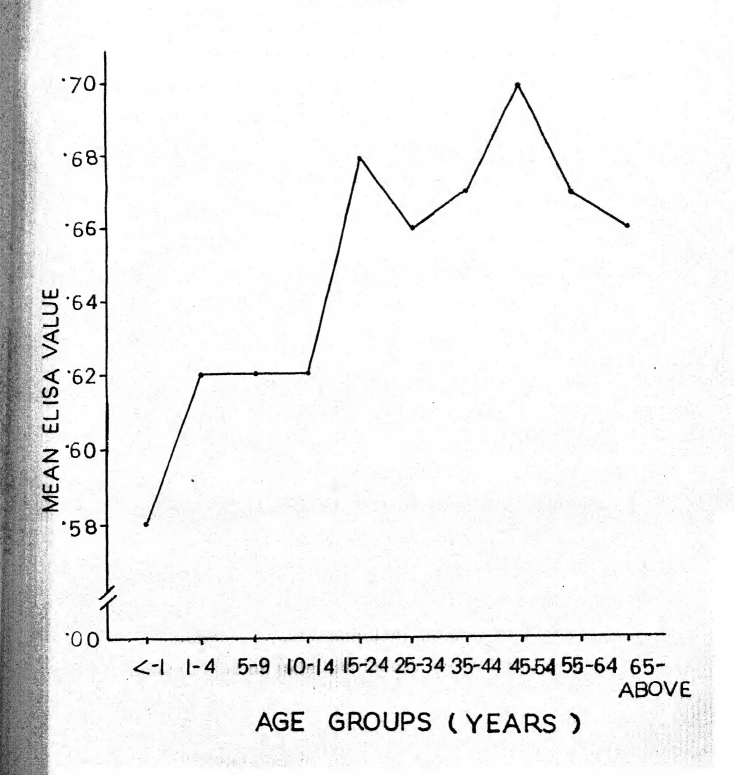
ages 2 - 10 years showing malaria parasite in their blood
films. The parasite rate was higher (2.16%) in female and
lower (1.59%) in males. Total parasite rate was 1.63 percent.
In the same age group sero-positivity was higher (15.63%)
in females and lower (13.23%) in males.

Distribution of ELISA value according to age.

				CONTRACTOR CONTRACTOR CATALOG			67.10		
		0 - 0.2 0.3 -	0.3 - 0.4	0.4 0.4 - 0.6 0.6-0.8 0.8-1.0 1.0-1.2 1.3-1.4	0.6-0.0	0.8-1.0	1.0-1.2	1.2-1.6	To Les
•									9.3
•	3	23	200	6.93	(3,88)	*:	(2,88)	(0.96)	0.62
•	\$	\$ 3.5	35	(9.09)	(3.53)	(1:52)	(0.50)	(0.50)	0.62
3	8		(8.78)	#	(4.47)	(100)	(1,60)	•	0.62
8	•	30.33	20.00	(19.69)	(15.09)	(S. 36)	(1:35)	(0.62)	0.66
1	:	# #	\$600	(38.46)	(22.36)	8.5	(3.25)	(1.22)	99.0
3	E	(10.17)	(17.51)	(38.40)	¥	(10.73)	(3.39)	(0.36)	0.67
8	8	100.000	10000	(10.00)	2 % 3	\$ \$ \$	3.5	(3.00)	6.3
8	3	(4.76)	33	(42.96)		(11.43)	(3.61)	•	0.67
•		3.60	02.60)	(40.72)	(3.66)	3.6	(3.86)	•	9.6
la	E	(17.86 (17.88	(36.98)	(24.61)	25.00 (3.00)	(60.05)	s.s	(0.53)	•

TABLE 15

GRAPH SHOWING MEAN ELISA VALUE BY AGE



# 6. ELISA Test :

#### 6.1 ELISA by Age :

showed that the mean value of ELISA increased with the increase in age upto the age of about 54 years (Table 18, Figure 16). The mean values of ELISA showed a decreasing trend in the alderly age group i.e. over 55 years. The individuals showing the ELISA value of 7% .40 E492 optical density with serum dilution 1 : 400 were taken as ELISA positive at this cut off point (Ray et al., 1983).

#### 6.2 ELISA by Sex :

There were 28.36 percent male individuals who showed  $\angle$  .40 0.D. ( $\mathbb{E}_{492}$ ) value and also 26.57 percent individual showed  $\mathbb{Z}$  .40 0.D. ( $\mathbb{E}_{492}$ ) value. The mean ELISA value was 0.65 0.D. in case of male individuals. There were 24.01 percent female individuals who showed  $\angle$  .4 0.D. ( $\mathbb{E}_{492}$ ) and also 21.05 percent individuals showed  $\mathbb{Z}$  .40 0.D. ( $\mathbb{E}_{492}$ ) value. The mean ELISA value was 0.64 in case of female individuals. There were less difference seen in mean ELISA value of male individuals.

1	W.	
-		
State and		

4	. *	įį		į	*   *	ğ	ı k	*	* ×
		•	•	•	•	•	•	•	1
	10.00		33.00	•	48.00	•	20.00	•	•
	13.33	2	40.00	#	36.66	•	20.00	•	•
	34.32	=	29.73	2	35.14	•	9+11	**	2:70
	24.93	8	22.39	2	40.30	8	20.90	M	1.49
	27.72	2	17.73	\$	37.97	5	26.58	•	•
	8.2	3	22.23	2	39.68	2	22.43	•	•
	19.01	2	21.70	2	35.65	*	22.64	**	16.0
	13.10	2	21.52	28	44.33	z	30.38	**	1.37
	20.68		20.69	3	18.29	•	10.34	•	•
		3	20.78	325	37.95	191	22.30	•	0.00

# 7. IIF Test Reaction :

# 7.1 III reaction by age :

Table 19 shows the reaction grading at 4+ with the test dilution giving a 2+ reaction used as the end point is ELISA positive individuals. Of the 722 ELISA positive individual tested with ITP, 132 (18.28%) have shown no reaction at 32 dilution end point or only a trace of fluorescence; 150 (20.70%) individuals showed a 1+ reaction which was only dimly fluorescent and very diffuse. There was 274 (37.95%) individuals samples have shown a 2+ fluorescent with the individual parasite quite diffuse, and 161 (22,23%) individual sample have shown a 3+ reaction which was brightly fluorescent but individual parasite was somewhat diffuse. There was 5 (0.69%) individuals who should a 4+ brilliantly fluorescent with the individuals parasite sharp and easily brought to focus (Hall et al. 1978). By IIP test, the sere-positivity was observed as 60.94% in MLISA positive proven individuals et 1 : 32 dilution oud point.

the bigher (30.37%) resetion grading was also chaeswad in age group 55 - 64 years, followed by 35 - 34 years age group. The lowest (0.11%) reaction grading was absented in 800 grading was

Metelbution of number of sers scording to reaction grading at 1:64 dilution end point by age. 0.38 8.00 5.75 \*\*\*\* \*0 25.00 16.46 26.26 23.62 11.19 14.56 8.73 10.39 10.34 13.63 \*\* 2 5 2 37.93 25.00 36.36 24.33 27.32 46.32 45.87 36.43 33.09 43.65 ‡ 263 11 25.76 16.46 27.50 20,00 19.18 10.25 18,87 3.99 33.54 40.34 2 34:52 21.52 34.14 12.00 13.51 23,13 34.68 24.53 21,31 19.33 274 712 1

Table 20 shows reaction grading at 1 : 64 dilution and point by age. There was 174 (24.10%) individuals who showed no reaction followed by a 1+ reaction in 186 (25.76%), 263 (36.43%) showed a 2+ reaction, 97 (13.43%) showed 3+ reaction and only 2 (0.28%) showed a 4+ reaction.

Out of 722 ELISA positive individuals, 372 (50.14%) showed a 2+ and above reaction. The positive reactivity in ELISA positive individuals was 50.14%. The higher (46.22%) positive reaction was observed in age group 45 - 54 years, followed by 55 - 64 years age groups and lower (24.32%) was observed in 10 - 14 year age group.

# 7.2 III reaction by sex :

Individuals according to their sex. Of the 494 males SLISA positive sers tested, there were 234 (57.92%) sers showed 2+ and above reaction grading at 1 : 32 dilution and point and 266 (42.00%) females. ELISA positive sers have also shows 2+ and above reaction at the same dilution and point.

	30 28			section Gradings				
1	1		1			*	*	**
	ė	×	ė	*	ě	×	ġ	×
		19.88	3	3.8	8	35.65	•	0.74
	2	2	3	*0**	2	23.27	~	3.
	2	20.70	34	37.56	161	22.30	•	198

Table 22 depicts the reaction grading at 64 dilution and point by sex. There was 196 (48.51%) male ELISA positive sexs which showed 2+ and above reaction grading, and 166 (52.20%) female sexs showed 2+ and above reaction. At 1:32 dilution, out of 318 (33.4%) female sexs, 206 showed 2+ and above reaction. There was 9.41 percent male and 12.56 percent female which dropped in 1:64 dilution and point.

Indirect Immunofluorescence antibody titres in ELISA positive

TABLE 23

sers using antigen from strains of cultured plasmodium felciparum by age.

No.of		IIF Antibody titre						
	a de la companya de l	256	120		- 33	14	8	
			-	•		•	•	
1-0			•	11	13	•	•	
5-9	33	•	•	20	17	•	•	
0 - M	57		•	17	17	•	•	
5 - 24	134	•		59	84	•		
6 - 24	150			66	103	•	•	
S - 48	136			66	77	•	•	
5 <b>-</b> 51	LOS	•		60	63	•		
5 - 64	70			49	50			
9.0	•			34	17	•	-	
otel	722			372	440			

(x<sup>2</sup>=5,209, 6,2,=0, 7 70,750), (x<sup>2</sup>=1,604, 6,2,=0, 7 70,97)

#### 7.3 IIF Ambibody titre by age :

Table 22 shows indirect immunofluorescence amtibody titres in ELISA positive sera, using antigen from stains of cultured plasmodium falciparum by age. By using in-vitro culture, plasmodium falciparum stain antigen, amtibody titre ranged from 1:8 to 1:256 dilution end point. In this test, starting dilution was 1:8 and significant titre were 1:32 and above by 95 confidence limit in relation to normals (Spencer et al., 1979; Mahajan et al., 1981). Therefore, an antibody level of 32 or more was considered as of sero-positivity. Amongst 722 ELISA positive individuals of all ages, 440 (60.96%) had positive titre (1:32) and 372 (51.53%) showed positive titre (1:64).

In relation to equ, sero-positivity rate was

Increasingly higher (9-14%) in age group 25 - 34 year

and lower (1.52%) in age group 1 - 4 years, and thereafter

sero-positivity rate showed a decreasing trend upto age

group (5+ (1.56%). The difference was statistically

not significant by titre (1 : 32). The difference was

also not algorithms. By titre 1 : 64 in relation to

TABLE 24

Indirect Immunofluorescent antibody titres in ELISA positive sera using antigen from stains of cultured Plasmodium falciparum by sex.

	No.of		IIF Amtibody titre				
	exemined	256	120	64	32	26	8
Rele	404	•		206	234		-
				(51.00)	(57.92)		
Penale	318	•	-	166	206	-	-
				(52.20)	(64.78)		

## 7.4 IIF entibody titre by sam :

and 1: 64 suspectively.

TABLE 25 Comparative results of slide positivity and sero-positivity by age.

Age group year)	Total No.of Sample examined	Slide		A THE PARTY OF THE	Positive by SLISA		ITY In BLISA		
				80. 8		1031755		1:6	:4 K
<u> </u>	Total Control	-	***	-	*		-		
1-4	108	•	3.70	20	18.52	13	12.04	11	10.19
8-9	190	1	0.50	33	16.66	17	8.59	20	10-10
10 - 14	179	3	1.68	37	20.67	17	9.50	17	9.50
15 - 24	201	6	1.87	134	41.74	84	26.17	59	10.30
15 <b>-</b> 34	246	11	4.47	158	64.23	102	41.46	66	26.63
15 - 44	<b>IN</b>	7	3.95	126	71.19	77	43.50	66	37.29
15 - 54	100	3	2.00	106	70.66	63	42.00	60	40.00
W - 44	100	•	3.01	79	75.24	50	47.63	49	46-66
			2.53	29	74.36	17	43.59	14	35.94
istel			2.63	722	47,50	440	60.94	362	50-14

mote a No individual face (1) year ego group was positive by

Catholic St. Shows samples to the state of t

TABLE 26

Comparative results of slide examination/ELISA/IIF Test by sex.

	Total .			19. of	Positiv	e by			
Sex	No.of Samples	81. exes	de <u>mation</u>	Constitution of the second second second	LISA	113	10 EL3	SA P	생빛의
		80.	*	No.	*	ъ.	K	Eo.	
Male	635	21	2.51	404	40.30	234	37.92	206	50.99
Perale	605	19	2,77	318	46.42	206	64.70	166	52.30
rotal	1520	40	2,63	722	43.50	440	60.94	372	51-58

# 7.6 Comparative results of slide examination/ELISA/IIP Test

ned sero-positivity by sex. Slide positivity rate was higher (2.77%) in females and lower (2.51%) in males. The percentage of sero-positivity was higher amongst male (40.30%) and lower amongst females (46.42%). In ELISA positive samples, IIP antibody titre was higher in males (57.92%) and (50.99%) makes at 1 : 32 and 1 : 66 dilution respectively, and was also higher in Semales (66.78%) individuals at 1 : 32 dilution and lower (52.20%) female individuals at 1 : 32 dilution and

ELISA velue end IIF reaction greding in slide positive individuals.

81.	ELIA velue	IIF reest	ion grading	51.	ELIEA Value	IIF Reaction Greding	
Code Bo.	ebeer- bence Unit	dilution	i : 64 dilution	Code Bo.	abser- bence Unit		1164 0110- 1100
<b>30</b> A	1.13	++	++	73	1.09	**	**
23 A	1.05	+++	+++	8 3	1.01	•	
25 A	1.14	**	**	10 3	1.07	4+	•
41 A	1.01	**	•	16 L	1.10	+++	**
50 A	1.05	++		17 L	1.13	+++	+++
52 A	1.16	++	+++	10 L	1.05	+++	++
56 A	1.46	+++	+++	19 L	1.13	+++	+++
67 A	1.00	**	**	21 L	1.33	**	++
25 B	1.15	+++	+++	22 L	1.23	**	**
20 B	1.00	•		23 L	0.99	+	+
22 B	1.13	***	+++	24 L	1.09	**	**
30 B	1.10	***	***	26 5	1,50	+++	**
35 B	1.21	**	•	32 N	0.94	++	++
36 B	1.35	**	**	3 .	1.00	++	+
79 Y	1.01	***	***	4 2	1.02	++	
22 N	1.20	*	•	10 9	1,16	+++	+++
			New States	11 7	1.15	***	+++
16 I	1.00	•		12 >	1.09	***	+++
	1.03	•	*	13 P	1.03	***	**
	1.24			17 >	1.10	+++	**

#### S. Sero-positivity in slide positive cases :

grading in slide positive individuals. Of the 40 individuals who were slide positive, all the forty were ELESA positive as well. They have shown ELESA value ranging from 0.99 to 1.46 optical density at 492 n.m. and in all of them. three to four fold rise of ELESA value (more than 0.39 optical density at 492 n.m. ELESA value) would be demonstrated. Of the forty confirmed malaria positive individuals, 36 showed II7 test positive reaction at 1 s 32 dilution and point and 31 showed positive reaction (2+ or more) at 1 s 64 dilution and point.

TABLE 18

Velidity of ELEMA test at above 39 absorbance unit.

RLBA results	<u>Slide reculta</u> Positive Negative	Total
legative		722 796
	1400	1520

Validity of IIF test (a) at out off titre of 32

IIF results	Post Edve	Magatira	Total
Positive	36	404	440
Negative	113.	270	202
Neal Control	40	682	723

TABLE 30

Validity of IIF test (b) at out off titre of 64.

IIF resules	Footsive Regults	Total.
Positive		972
Wegetive		
lotal .	• •	723

## 9. Velidity of ELISA test :

of the test was calculated at the 3.39 optical density (E<sub>492</sub>) as out off point. It was found that specificity of the test in low transmission study area, was 53.92%. Test was very sensitive at the '7.40 optical density (E<sub>492</sub>) out off point and showed 100 percent sensitivity in slide positive individuals.

#### 10. Velidity of IIF test :

In the area, the specificity of the test at cut off titre of 1 : 32 was 40.76%, at 1 : 64 it was 50.00 percent.

The sensitivity of the test at the cut off titre of 1 : 32 was 90.00 percent, at 1 : 64 it was 77.50 percent. Whereas at cut off titre of 1 : 32 though the sensitivity was good (90.00%), the specificity was lawer (40.76%). At cut off titre of 1 : 64 though the specificity was slightly better but the sensitivity was poor (77.50%).

Hence, a cut off titre of 1 : 32 appeared to be best suited for studying the sero-epidemiology end was used in the study.

\*\*\*\*\*

DISCUSSION

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#### DESCRIBE THE

Malaria continues to be an important health problem in most of the countries of South gast Asia Region and requires a flowable epidemiological approach with available resources. However, the success in integrated malaria control could be seriously impeded without a sound knowledge of local epidemiology of malaria. Since during past three decades, socio-economic status as well as the habitats and eco-systems of malaria vectors and parasite, deforestation, flooding, irrigation, the green revolution, vectorial efficiency of anophilines and their sibling species and uncontrolled population movements have also had a significant impact on malaria epidemiology (T.R.S. 547).

The serology of malaria has a long and verted bistory. Only in the last two deceds, now escalagic techniques such as the indicect fluorescent entitledy test (III) and Emayor linked immensurbent usery (NiIA) have been explored for use in solving problems associated with epidemiology, specialist and diagnosis.

## 1. Population under study :

The study under discussion was conducted in reral area of Chirpson, Themsi (U.P.). A population survey for

Company of Post St

slide and seropositivity for meleria was carried out in the area. The studied population consisted of 1520 individuals from 269 families out of 290 families. An ettempt was made to include all ages and both sense in the study to know the serological profile of the population. The study area was known hypo-endemic (API 2.6) for malaria. Hence it was decided to have 20 percent sampling of the population to look for correlation between serological and slide positivity.

Lobel of al (1976) is sero-epidemiological investigation of malaria in Guyana, where malaria control was in maintenance phase included 20 percent of the estimated population of survey sectors. Name at al (1979) took a small sample. They examined only 60 and 50 households in first and second surveys. The total population of surveyed area was 30,033.

1987, wince it was post of transmission secondary of transmission secondary of transmission secondary of transmission secondary and transmission of the transmission of both the species viscential control for the transmission of both the species viscential control for the transmission of both the species viscential control for the posts months. It was due to be posts months. It was due to

## 1.1 Male Pemale ratio :

The male and female ratio of sample in our study was 1000: 818 as compared to 1000: 845 for the population of area (Census, 1981). The male female ratio was less as compared to Census 1981. The young adults male and females contributed 30.92% and 31.37% respectively. The prediatric population sampled in the study was 31.72 percent. Male accounted for 18.03 percent and female 13.69 percent. However, only 1 (0.07%) individual was in age group \_ 1 year sampled during survey. This disparity was due to non co-operation of perents of children and adults were not available during day time and due to unavoidable circumstances, could not be contacted in the evening in that particular village.

There was poor co-operation from adult females due to practice of purdha. They refused to come out from the house for interrogation and investigations. But this was not expected to effect the results as sex differences do not produce variation in serological studies for malaria (Lobel et al. 1976).

## 2. Bio-eccial Characteristics of Population :

#### 2-5 400 \*-

The present study revealed that the alide positivity rate (SPR) for malaria varied with app. The bigher (1.48%) slide positivity rate was observed enought individuals of 25 years and above. It was found lower (0.50%) emongst individuals belonging to age group 5-9 years. Slide positivity was 3.65 percent in the 1-4 years age group which indicates that fresh transmission is occurring in this area. He comment on age group \( \subseteq 1 \) year could be made due to non-availability of samples from this age group. There was no significant (P \( \subseteq 0.25 \)) statistical difference in slide positivity rates between age group 1-4 years and 15-64 years.

Choudhery gi gl (1986) cerried out a study and classified the population on the basis of the clinical history of malaria observed that all age groups were affected by the disease but that there was a progressive increase of malaria attacks to 16-25 years of age, where the rates reached the maximum level.

Pattemayak at al (1978) while attempting to amelyse the dynamics of re-establishment of salaria encemicity, found that the malaria incidence in the age group 5-14 years was quite high oven at initial stages of salaria epidemics in comparison to other age groups. The prevalence of malaria among different age groups is subject to wide veriations. According to Mandonaid (1987) in the salaria provalence to runnis note or lass evenly distributed many different age groups during malaria epidemics. The value of malaria testage groups during malaria epidemics. The

used extensively for identification and specification of endemicity in malarious area (Pumpana, 1969). The dynamics of these indices in areas subject to anti-melarial measures served as a basis for assessment of these measures (Bruce-Chavatt, 1985).

Verma <u>et al</u> (1983) observed overall slide positivity rate of 3.42%, without any significant difference for various age groups. This corroborates well with our study.

The higher (7.8%) slide positivity was observed by Mamtani at al (1982). Uprety at al (1982) observed 6.8% slide positivity rate in afebrile healthy children of 2-9 years of age, although the positivity was higher (45.5%) in febrile individuals.

percent was seen in 1-4 years ago group and higher (75-24%) in 55-64 years & shows ago groups. Ago has been an convenient variable in interpreting the results of serological technique. This can tall whether transmission occuring is fresh or it is previous experience of malarisally ago groups are expected to be involved where fresh transmission of malaris is occuring. The immunity to malaria sions as the age of individual increases with consequent reduction in paramite rate (Descults 2% Malaris 1968). The Salarish has conducted secological studies in accountly be accounted by a salarish by a salarish areas of the country.

during 1970s, and it was observed that the population below 5 years of age had hardly any malarial emperience. It was only higher age group who showed high titres.

This correlated very well with our observations that the sero-positivity was significantly low in the age group 1-14 years when compared with 15-44 years and 45-64 years age group (P \( \sum 0.001 \)).

The overall sero-positivity was 45.7 percent.

Out of the total ELISA positive individuals, 60.94%

individuals were positive at 1.32 dilution and point

by indirect immunofluorescence (IIF) test and 50.14

percent at 1:64 dilution were also positive by IIF test.

In the literature, the data on comparative studies of RLISA and LIF are extremely scarce, our findings are in accordance with Collins gt gt (1972) who observed percentage positive to <u>Pafalciparum</u> antiqua ranging from 47.2% to 86.0% and for <u>Plasmodium folciparum</u> or <u>P. malariae</u> antiqua percentage positive ranging from 60.0 to 80.0% and for <u>P. malariae</u> from 52.8 to 80.4%.

Colline of al (1972) also reported 78.4 percent sero-positivity for younger groups and \$2.5 percent for older groups. It is appeared that in younger upo group the III test falled to detect a sumber of those with potent person person person person person person to the person of the

In our study, 100 percent of malaria cases could be detected by a P-falciparum ELISA test and 90.00 percent at 1:32 dilution and 77.50% at 1:64 dilution by IIP test. This corroborates well with Srivestava at al (1983) who observed 100 percent sero-positivity of malaria patients (Group II), 28.78 percent of patients varied origin (Group II), 58.75 percent of random hospital patients (Group III) and 15.48 percent of the normal healthy subjects (Group IV). Ray at al (1983) obtained 85.1 percent sero-positivity in malaria cases by a P-falciparum ELISA test. Agarwal at al (1981) obtained 90.7 percent sero-positivity by using P-cynomulgiantigen and 79.9 percent sero-positivity rate by IIP test, using P-knowlesi antigen.

#### 2.2 <u>Sex</u> 1-

Penale dominated the scene in present study as alide positivity was higher (2.77%) then makes (2.51%). The difference was not significant (9 70.75). On the contrary, Srivestave at al (1975) observed higher slide positivity rate for makes (4.60%) as compared to females (2.29%). Makes sleep outdoor more than females, thus regulating in a frequent man-masquitoes contact.

Deljaer <u>et al</u> (1986) in Mayurbhamj District (Osiese), did not observed significant difference in slide positivity rates in males (12.5%) and females (10.6%). It was not clear whether it was due to higher exposure of adults males to malarie or due to influence of local socio-economic status, ethnic groups, or the attitude of parents especially mothers towards male and female children regarding treatment and ignorance about evailability of free services in the village. There was no relationship of sex to species of malaria parasites and no significant difference between infection rates of males and females with any species of parasite (Sweet, 1933).

Sero-positivity rate was higher in males than female and the difference was statistically significant (P \( \int\_0.25 \)). Similar observations were made by Collins at al (1971) in a study carried out in Ethiopia, found seropositivity 36.7 percent and 4.3 percent at low and high attitude respectively. The seropositivity was higher among males than females.

## 2.3 Religion & Caste !-

In our study, majority (48.60%) belonged to beckward casts, followed by scheduled casts (35.72%). 
Upper casts (14.8%) and Muslims (0.86%). Nowever, the slide positivity rate was highest (7.69%) for muslims, whereas for scheduled and backwards it was found to be 3.68 and 1.89 percent respectively. This difference was not significant (9 7 0.10).

Srivestave et el (1975), in the same district found more cases amongst Hindus which largely reflect the population composition during recent years.

#### 2.4 Marital Status :-

The present study revealed high (3.19%) slide positivity rate in married and lower (1.93%) in unmarried. The difference was significant (P \( \sum\_{0.01} \)). The meropositivity rate was highest (65.80%) in widow/widower/divorces, followed by (62.22%) in married and lowest (22.77%) in unmarried individuals. The difference was significant(P \( \sum\_{0.001} \)). It is due to the fact that married individuals belong to higher age group.

#### 2.5 Literacy Status :-

our study revealed that meleria is more common amongst illiterates. Slide positivity rates amongst illiterates and literates were 4.29 percent and 2.35 percent sespectively, which was statistically significant (P \( \subseteq 0.025 \)). On the contrary, Verma at al (1983) did not observed any mignificant association of slide positivity rate with literacy status.

The sero-positivity was \$0.76 percent in Aliterate. It was found that ease-positivity into declined with increasing literacy status.

#### 2.6 Occupation :-

association between slide positivity of malaria and various occupations. Association between sero-positivity and various occupations was also significant (P \( \) 0.001). The slide positivity rate was higher (11.11%) in individuals engaged in service in thermal power project, railways etc. and declined amongst farmers (2.94%). housewives (3.15%), followed by labour (2.04%). It was lower amongst student (1.65%) and children (1.66%).

enses occur among various entegories of Agricultural
labour (Pattanayak, 1981). The rest of cases occur in
unben and other areas of the country (Sharma, 1986;
Kondrashin & Dimit, 1985). The misk to acquire malaria
is higher among mobile workers and among those emposed
to manguite hites in open air on account of their
accupational requirements (Kondrashin, 1986). SPR and
slide falsiparum rate (SPR) in particular, was higher in
labour force engaged in tem plantations, in forcet account
membro cutting in jumgles, as compared with same index
among local inhabitants of meighbouring area (Mational
Malaria Eradication Programme, 1984). Construction
workers at development projects, fishermen, coal miners
and labour employed in number of thermal power projects

as well as reliveys in the peripherel part of the country show a relatively new pattern of labour movement and had shown higher SPR and SPR. There was explosive malarie situation with evidence of chloroquine resistant P. Esleipszum (Say, 1984; Raj Copalan, 1984; Panicker et al. 1984; Panicker & Raj Copalan, 1986).

Comparative sero-epidemiological studies among migrant workers and the sedentary population residing around Sathanam Reservoir in Tamil Nadu revealed that the Sormer had a higher positivity rate as compared with the latter (Nyma & Ramesh, 1980).

#### 2.7 Secial Class :-

our study revealed a higher (4.35%) slide

positivity rate amongst individuals from Social Class V

to those from Social Class IV (2.02%) and Social Class III

(2.41%). No individual was found positive amongst Social

Class I and Social Class II. The difference was significant

(p '7 0.05). It was due to low socio-economic status,
individuals were living in ill-ventillated, ill-lighted

and unhygianic houses surrounded by various types of

water collections. Poor people usually live in huts/
hutchs houses and keep cattle inside their residences.

thus resulting in mosquite meetling places with them.

Verms of al (1983) has reported higher SPR (3.61%) for
those belonging to social class V as squinst about 2% for
Social Class IV.

The sero-positivity rate was higher (54.48%) in Social Class V, followed by Social Class IV (46.10%) and Social Class II (40.74%). Difference was significant (P 78.010). Malaria, though common in all groups of society, has been significantly increasing among economically backward classes, inhabiting areas with difficult accessibility on the periphery, and where melaria eradication was never achieved (Rey, 1979; Kondrashin, 1983).

#### 2.8 Over-crowding :-

alide positivity rates observed by us were

2.93 percent and 2.22 percent in individuals residing in

over-crowded and uncrowded conditions respectively. The

difference was not significant (P \( \) 0.25). The sero
positivity rate for individuals residing in erouded and

uncrowded conditions were 49.99% and 45.41% respectively.

This difference was insignificant (P \( \) 0.01).

governed by the presence of parasite vector and subtable savironmental conditions in the community. Its distribution weries from village to village, and town to town and over from ward to ward in the same community depending on malariogenic conditions. Mondrashin & Orloy (1985) descred positive correlation between Extins incidence and population density as such the most intensive focal of Extras are altered usually in over-populated plain area.

3. Slide positivity and seropositivity in relation with Clinical Manifestations:

#### 3.1 Manatomegaly :-

In our study, slide positivity rate was higher (17.31%) in hepatomegalic individuals and lower (2.11%) in individuals without hepatomegaly. Repatomegaly was significant in slide positive individuals (P \( \) 0.001). The sero-positivity rate was higher (94.23%) in individuals with hepatomegaly and lower (0.20%) in individual without hepatomegaly. The difference observed was significant (P \( \) 0.001). Descrits & J.J. Seave (1965) in a study of immunity to malaria in protected and unprotected groups showed the liver enlargement rates, for all age group were lower than spleen rate but with the advance of age there was a decrease in liver enlargement rates. Liver rates were strikingly decreased of the protected population in contrast to unprotected population.

In congenital malaria, hepatomogaly and joundice with hemolytic ensemie is common in an infent. The diagnosis is confirmed only by detection of the malarial parasite in the paripheral blood of the infents (Thompson et al. 1976).

## 3.8 Balancasair e

Out of 1500 individuals, 144 (9.47%) had splanamously and showed 12.5 percent alike positivity.

Purther analysis of these 144 individuals with splenomegaly 92.36 percent showed sero-positivity and rest were sero megative. Slide positivity was 1.60 percent in individuals without splenomegaly. The difference was statistically significant (P \( \int 0.001).

Thomas et al (1981) conducted a sero-epidemiological study on aborigine children in Orang Aseli Malaysia, revealed that the falciparum antibody prevalence rate was 84.6% as against to spleen rate (81.8%) and parasite rate (43.3%). There was positive correlation between sero-epidemiological study and spleen rate, particularly in the age group upto 9 years old. Splenomegaly is a good clinical manifestation for diagnosis of malaria during epidemics and in hyper-endemic areas as it gives on the spot results but it is of no value in low endemic areas where it does not depict the true prevalence, nor it is useful in monitoring the progress of meleric control programme. All patients with splenomegaly do not have melaria and all patients of malaria do not have splenomagaly. In view of the fact that the population in this zeral community do take prompt presumptive treatment which provents spleen from becoming enlarged, and that there is mogligible difference in individuals with and without Parasitaonia. However, higher sero-positivity rate was found in individuals with spleasmoguly, consequent upon a sustained malaria challengeIt was unlikely that malaria was the actiological factor in the splanemegaly of these individuals. These results therefore confirm that splane enlargement is an unreliable method for epidemiological assessment of malaria when, as at present, widespread use of unti-malarials is provalent. Vander Kany also obtained similar results in an epidemiological study carried out in Surinam in 1972-74. Nameani gt al (1979) also obtained similar results in a serological survey for malaria in a rural community of Delhi.

#### 3.3 Annels :-

was the second

In the study, slide positivity rate was more in ammenic individuals, but all the individuals were serologically positive. This is due to the fact that they have suffered from malaria in the post and there may be other causes of america in the population studied. All of the 40 individuals with slide proven parasitaemia did not show america at the time of survey. It was clear from the results of slide positivity that america is not a constant manifestation in recent infections, whereas, 2.45 percent slide positivity in non-america individuals indicate that america is common in chronic malarial infections as a remote manifestation.

#### 4. Past Mistory :

#### 4.1 Past history of fever :-

There were 47.50% individuals with past history of fever and \$2.50% individual without past history of fever. The slide positivity rate was higher (3.60%) in individuals with past history of fever and lower (1.75%) in individuals without past history of fever. The difference was statistically significant (P \( \subseteq 0.01)\). The sere-positivity rate was higher (92.94%) in individuals with past history of fever and it was lower (6.16%) in individuals without such history. The difference was statistically significant (P \( \subseteq 0.001)\).

rever was classified as typical when it was intermittent and associated with chills and rigors; and atypical when it was either continuous or remittent without chills and rigors. Sharma gt gt (1985) observed body temperature in malaria patients ranging from 37.3°C to 41°C. Yever was more agate in Taleiparum than Yivex malaria.

The world therefore he had to complain that forest seems and reliable symptom of melania.

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Exercised 25 M (1979) the descript the seem pleasurement

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## 4.2 Past history of treatment (Presumptive/Radical) :-

The slide positivity rate was higher (2.95%) in individuals with past history of presumptive treatment and lower (2.42%) in individuals without past history of treatment. The difference was not significant (9 70.75). The sero-positivity rate was higher (97.87%) in individuals with past history of treatment taken and lower (1.43%) in individuals without past history of treatment taken. The difference was statistically significant (P / 0.001). Serology in general, confirmed and extended results of slide exemination and it was successful where slide examination failed, in detecting persons with melaria contact and possibly with sub-patent parasiteemia. Some of these reactions may have represented residual antihody from cured infections. Following enti-malarial treatment antibody titre declined. Schizonticidal drugs or natural immune responses of individuals may decrease antibody titres from significant levels. Such infections in individuals may exist for almost 1 year but this is only a minimal time (Mornstein at al. 1983).

## 5. <u>Parente Late</u> !

In the study, the original parameter rate was

L-03 percent. It was higher (2-169) in females and lower

(1-269) in males. In the age group 2-10 years, the higher

exce-positivity (15-699) was choosing in males as compared

to females (15-250): Studies (16-610) was reported by

Mamtani et al (1979) in a rural community of Delhi, who observed paresite rate 3.22 percent. Verma et al (1983) revealed an overall parasite rate of 3.42 percent which was higher (4.60%) for males and lower (2.29%) for females. The difference was not significent. On account of quite scanty work in recent years on the type of subject considered, the results of present study are difficult to be interpreted widely. However, in a study in mesoendemic area, Uprety et al (1982) obtained 6.8% slide positivity rate in afabrile healthy children of 2-9 years. although the positivity was higher (45.5%) in Sebrile cous. Slide exemination can only indicate the presence and absence of patent parasituania at the time of examination. It does not indicate individual malaria experience (Kagan, 1972). Absence of patent parasitamnia can be misleading since patency is influenced by immune status of individuals and use of enti-malazials.

A survey carried out in the Combin (Herreron,
ELLoon & Hell, 1948) showed good concertions between the
parasite rate and 2-A. tests in the rural area when the
transmission was at high level. Tellor 25 25 (1980)
conducted a study An Mark Africa Serezus, making Hills 
testsalone, found that many ruray infants were Siles negative
even though they had provious proven perceitemate. This
may be don to those going infants are not sufficiently
mature to meant an affective honoral suspense and forman-

#### 6. BLIBA Test :

## 6.1 Yelidity of ELIEA toot :-

Emsyme linked immunosorbent assay has been applied in the diagnosis of many infectious diseases (voller et al. 1976). The present study reports evaluation of micro-ELISA in maleria and confirms the finding of May et al (1963). That in vitro cultured P. felciparum is an excellent source of entiren for this test to detect antibodies. Between the three untigen batches the P. felciparum strain had undergone over 100 more sub-cultures. Perssite did not show any changes so far as entiqualcity is concerned with regard to micro-ELISA test. It was also obvious that with higher parasitagmia of the culture, the yield of antigem was more. The date on replicate testing of the test and reference sore within the same batch as well as with three different batches of antigons showed consistent results indicating the usefulness of the test. In this report, at a serum dilution of 1:400, \$3.4 percent of normal individuals showed reaction upto 0.4 whereas the rest showed reading between 0.4 - 0.6 0.D. In contrast, Spencer at al labelled a reading of 0.3 at dilution of 1:40 as negative and so positive reaction was noted assesst sormal individuals at 1:00. In a study on a sormal healthy Indian population (Mahajan <u>et al</u>, 1991), maing Autus P. Paleiparm antigen, non-specificity was found to be 6.06 percent which is comparable to our results. However.

in this study the results were read visually 1 : 100 was considered as the cut off point.

In our study, at a serum dilution of 1 : 400, \$2.50 percent of individuals showed reaction upto 0 - 0.4 0.5.  $(E_{492})$ , 24.01 percent showed 0.4 - 0.6 0.5. at  $(E_{492})$  reaction, 14.81 percent individual showed 0.6 - 0.8 0.5.  $(E_{492})$ , 6.05 percent showed 0.5 - 1.0 0.5.  $(E_{492})$  and rest other showed upto 1.4 0.5.  $(E_{492})$ . All the slide positive individual showed sore than 1.0, 0.5.  $(E_{492})$  reaction.

The sensitivity and specificity were calculated at cut off point ( 7 .40). The 100 percent consitivity of the test in individuals with patent Parison parasiteemia reported here confirmed earlier studies. There was no difference in sero-positivity in case of first attack and more than one attack showing that this test can detect very early entibodies (Nebejen et al. 1981). Ray et al (1983) also observed 100 percent sero-positivity in all the 11 cases of <u>P.folciperum</u> infection from Maryana State. Similar observation were made by Voller and colleagues who found that this test was positive in 19 out of 20 Temmenian sera and in all the 41 Frances patients who were parasitologically positive for meleria. Dutte at al (1982) reported, using P. falciparum entigen, sere at a single dilution (1 , 200) from 143 maleria patients (Group I), 70 patients of varied origin (Group II). 75 random hospital patients (Group III)

and 75 normal healthy subjects (Group IV) were tested at a cut off point equivalent to 95% confidence limit of normal subjects (Group IV), 100.0, 6.6 and 12% cases respectively of Group I to IV gave positive ELISA reaction. Srivestave et al found that using P.falciparum antique, the ELISA test at 99 percent confidence limit gave 99.3 percent positive results among 143 malaria patients while none of the 70 patients of pyremia and 75 random hospital patients gave positive reaction. These findings confirm previous reports on other serological tests, (Mathews et al., 1975; Ray et al., 1983; Agarwal et al., 1968 and Wilson et al., 1975). Chandmani et al. (1981), Ragan et al. (1989), Mahajan et al. (1981, 1982) observed lower sensitivity as compared to present study.

The specificity was observed SJ-92 percent at out off tiere ( 7/.40) at which sensitivity and specificity were more acceptable. Similar observation was made by Ray gg gg (1983) who compared EMA, EIF and ELISA, and did not reveal eightforward EMA, EIF and SLUSA, and wilson gg gg and differences observed in the three tests suggest that mutibodies detected may comprise similar and dissimilar classes. Namejan gg gg found SLUSA to be far superior to IMA and EIF is newto malaria infection uning g. housing antique.

## 6.2 KLISA Test by ege and sex :-

In our study, ELISA value was increased with the increase in age upto the age of about 54 years and the mean ELISA value showed a decreasing trend in the elderly age group i.e. over 55 years at out off point 2 40 0.D. (8,40) with serum dilution 1 : 400. Our finding corroborate with Voller et al (1980), who observed in a longitudinal study of malarie in West Africa Sevense, in unprotected and protected population after one year of protection that showed ELISA values increased with age and reached a plateau by age 19 - 28 years in unprotected population. In the protected population, the ELISA values were significantly lower in age groups 1-4 to 9-18 years, but there was less difference in the older age groups at cut off point ( 7 0,2) with serum dilution 1 : 100. High SLISA value correlates well with degree of exposure. Halaria control activities result in low ELISA value. ELISA may give megative results in infants with provem parasitaemia (Voller et al. 1980).

The state of the control of the cont

Ray 21 al. 1983e, Dutte 21 al. 1982, 1984; Spencor 21 al. 1979, 1981; Voller 21 al. 1974 a.b. 1975, 1978, 1980).

Spencer gt al (1979 a) used in vitro culture of 
¿. falciparum entique for micro-ELISA. Positive ( "7 80)

ELISA antibody responses were found in persons with 
parasitesmin. In all the semi-immune individuals titre 
were 780, and reciprocal titre rose rapidly to levels 
72560 by 2nd to 9th day of patent parasitesmia and 
gradually decayed after curative therapy. In non-immune 
individuals titres were lower than in semi-immune 
individuals. However, positive titres do appear rapidly 
with patent parasitesmis. In another study (1979 b) they 
observed discordance between EIF & ELISA in 23% samples 
from Vietnam and 29.4% from Houndans. ELISA was magative 
in considerable number of parasitologically positive cases. 
Edison at al (1979) observed higher ELISA values in 
unprotected population in comparison to protected population.

## 7. Indirect Insunofluorescent Test :

## 7.1 IIP by Age & Bez t-

drawn between LIF & ELINA. However, higher (30.37%) reaction grading was observed in age group 55-66 years. followed by 25-34 age group. The lower (8.11%) reaction grading was observed in age group 10-14 years individuals at 1:32 dilution end point. Similarly, higher (46,22%) positive reaction was observed in age group 45-54 years. followed by 55-64 age groups and lower (24.32%) was observed in 10-14 years age group individuals at 1:64 dilution and point. The reaction grading (2+ and above) for males, was 57.92% and 48.51% at 1:32 dilution and 1:64 dilution respectively. For females, it was 46.78 percent and 52.20 percent et 1:32 dilution and 1:64 dilution and point respectively. In relation to age sero-positivity rate was increasingly higher (9.14%) in age group 25-34 years and lower (1.52%) in age group 1-6 years and thereafter sero-positivity rate showed a decreasing trend upto age group 65+ (1.96%).

Station to degree of translation, the labeletance of the Course of the C

indicated that the rate of increase in antibody is rapid in young children but slows down in adolescence and adult life.

collins et al (1971) in a study at Sthiopia.

observed 34.7 percent and 4.3 percent IFA positivity at

low and high sititude. This corresponded with other

maleriametric indices. The positivity was higher among

males then females. While studying antibody response in

persons previously exposed to maleria. Stude Charatt et al

(1972) concluded that about 50 percent showed a positive

response at low titre equinat P. felcipsum and P. Tital.

There was little evidence of persistence of maleria

infection in this group.

observed 0.8 percent and 31.7 percent positivity in population under 15 years and over 15 years tempertively. They suggested that positive responses were more likely to be associated with old or imported cases than with current local transmission. Srivesture 32 91 (1989) observed its high disquestic value since 90 percent of alide positive malarie patients carrying 2. falcingsum or 2. Nivel could be disquessed. Purthernore, positivity characteristic relies sold as senior hospital patients reflected a low degree of Selected positivity due to part oxperience of malarie facetains ments these cases.

## 7.2 Validity of III Test :-

Out of 722 total ELISA positive individuals examined by IIF, antibodies were detectable in 47.5 percent individuals from study area during transmission season. Some of them were not having malaria at the time of survey. In many of them detectable antibodies might have been due to previous experience to malaria. It was honce considered desirable to find out a cut off titre at which the diagnosis of malaria could be made with reasonable sensitivity and specificity.

The sensitivity and specificity were calculated at cut off titres of 32 and 64. The test was very sensitive at these titres and specificity range from 40.76 percent to 50.0 percent. Whereas, at cut off titre 1:32, though the sensitivity was good (90.00%), the specificity was lower (40.76%). At cut off titre 1:64 though the specificity was slightly better but the sensitivity was poor (77.50%). Taking this into consideration, cut off titre of 32 was taken for study at which sensitivity and specificity both were most acceptable and this was 90.00 percent and 40.74 percent respectively. In previous studies Aparwal gt gl (1901, 1902), may gl gl (1902, 1903) also found cut off titre of 32 to be most acceptable. Similar specificity was also observed by May gt gl (1903).

Higher specificity 67.9 percent was observed as compared to present study using same entigen (Ray gt al. 1982). Whereas sensitivity was much lower in evaluation study conducted by Colline gt al (1981) and Warren gt al. (1975).

Stuce Chewatt <u>et al</u> (1972) concluded that about SO percent showed a positive response at low titre against <u>P. falciparum</u> and <u>P. vivez</u>.

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CONCLUSIONS 

#### General room

The study was conducted in sural areas of Jhansi (hypo-endemic for malaria). One thousand five hundred twenty samples were collected during transmission posied (September - October 1987). Samples were examined for slide as well as sero-positivity. Also 722 ELISA positive samples, were further analysed for sero-positivity using IIF technique. The observations of the study have led to following conclusions:

- 1. There was no significant (P \( \sum\_{0.25} \) difference in slide positivity rate (SPR) among different age groups but sexo-positivity rate was significantly low (P \( \sum\_{0.001} \)) in those aged \( \sum\_{1} 16 \) years in comparison to over 15 years age group.
- 2. We significant (P 70-10. P 70-25) difference was observed in SPR and sero-positivity rate in relation to religion and oaste.
- s. Significant (P (\_ 0.0), P (\_ 0.001) difference was observed in both the rates between unacerted individuals and national individuals.
- o. Significant to 7 0.000, P (\_ 0.001) difference was absorbed to 500 and separately between

illiterates and literates as SPR declined with improvement in literacy status.

- 5. No significant difference (? ( 0.5) was observed in SPR for various occupations except individuals classified as other groups. While comparing adults and children, there was significant difference (? ( 0.001) observed in sero-positivity rate.
- 6. There was no significant difference (P 7 0.05)
  observed in SPR in relation to various social class
  but significant difference (P 7 0.01) observed in
  seco-positivity rate between Social Class II & V.
- 7. SPR was not significant (P \( \\_ 0.25 \) in relation with over-exceeding but sero-positivity rate was significantly higher (P \( \\_ 0.01 \)) in individuals residing in over-crowded duellings and heaping cattles within duellings.
- observed in SPR and Sero-positivity rate between individuals with hepatomogaly and without hepatomogaly.
- observed to the and sere-peritrity between individuals with optomic party and attended optomic party.
- 10. OPE and Description of Control Williams Research to the Control of Contro

- 11. We significant difference (P 7 0.75) was observed in SPR between individuals with past history of treatment and without past history of treatment but there was significant difference (P 4 0.001) in sere-positivity rates.
- 12. Significant difference (P (\_ 0.001) was observed in SPR between individuals with high temperature and without temperature.
- ineignificant among males and females.
- 14. The sero-positivity rate correlated positively with slide positivity rate.
- 15. The SLIBA values were markedly lower in age groups

  1 4 to 5 54 years, but there was less difference
  in older age groups. The SLIBA values increased with

  age and reached a plateau by age of 56 years, in

  protected population.
- 16. ELISA was found to be highly sensitive (100.0%) and moderately specific (53.92%) test.
- 17. IN was found to be semplating (SOLOGI) and medicintely specially (SOLOGI) test

18. Multiple serological tests should be performed for diagnosis of maleria. Rising antibody titre (ELISA) and raised IIP antibody levels alongwith any other positive test, give very strong evidence of maleria, but this needs further evaluation in an area with high incidence of maleria.

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#### APPENDIX - I

# SERO-EPIDEMIOLOGY OF MALARIA IN RURAL POPULATION

# PARTLY SCHOOLS

Village Code	30.	******* House No. ********
Name of Head of Pamily		**********
Beligion		Hindu/Nuslim/Other (specify)
Caste		Scheduled/Sachward/Syper
Main family occupation		*******
Type of family		Joint / Single
Total Mo. of family members		**********
Total monthly income (everage)		**
Per capita monthly income		Be
Social Class		1/11/111/14/4
No. of living rooms in the family		*******
No. of living units		******************
	•	Present / Absent
PARILY CONCESTION 1		
51. Hemo Age Sex will No. Hemo Age Sex will No.		ory Martiel Occu- acy Status pation Income the

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#### APPENDIX - II

#### SENO-EDIDENICLOST OF MALARIA IN RUBAL POPULATION

## INDIVIDUAL SCHOOLS

		COCH NO. 1
J. Washington		Date
Name of individual		***********
Age		***********
		Nale / Female
Village & Pamily No.		********
Sleeping hebits		Day / Hight
Past history of fever		******************
. If present, when ?		
. Deretion		*********************
· Competature		***********
·		Present / Absent
. Hepatomegaly		Present / Absent
. Splenomegaly		Present / Abrent
. Treatment II token	•	Presumptive only / Presumptive + Radical / None
. Sero-positivity		
1.1.7.		Present / Absent
ALL		Present / Absent
	*	Transmission period /
A Property of Paris 19		Non-transmission posted.
FIELD MOTES IF ANY :		

Signature of Investigator.

### APPENDIX - III

## Carbonate Buffer 9.6 pH:

- (a) 8.4 gm MeHoO, in 100 ml distilled water (Stock Solution-A)
- (b) 10.6 gm Mag CO2 in 100 ml distilled water (Stock Solution N).

100 ml of Stock Solution-A + 18.2 ml of Stock Solution-B and dilute to one litre of distilled water.

## Phosphate Buffer Solution pH 7.2:

0.5 M Ma\_MPO4 (35.632 gm/litre) Stock Solution-A C.5 M Ma M\_ PO4(31.200 gm/litre) Stock Solution-B

A + B dissolved in 200 ml of distilled water (Stock Solution)
40 ml of Stock Solution + 100 ml of 85% Aquous Heel.
Dilute to 1 litse with distilled water.

# Phosphate Buffer Solution Tween-20 (PBSIT)

1000 ml PBS with distilled water + 5 ml Tweep-20.

# Citrate Buffer pH 5-0 :

Stock A 3.84 gm of citzle acid in 200 ml.

Stock 2 5.68 gm of Me, MPO, in 200 ml.

35 ml Stock A + 25 ml. Stock B + 50 ml distilled vater is equal to 100 ml (Stock Solution).

\*\*\*\*\*